DESCRIPTION

METHOD FOR PREVENTING ELUTION OF NICKEL FROM WETTED INSTRUMENT OF COPPER ALLOY, PROTECTIVE FILM FORMATION AGENT FOR PREVENTING ELUTION OF NICKEL AND DETERGENT FOR PREVENTING ELUTION OF NICKEL

Technical Field:

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[0001] The present invention relates to a method for preventing elution of nickel from a wetted instrument of copper alloy, a protective film formation agent for preventing the elution of nickel and a detergent for preventing the elution of nickel.

Background Art:

[0002]Generally, valves for city water, feedwater or hot water, pipe joints, strainers, water faucet clasps, pump supplies materials, water meters, water purifiers, water feeders, hot water feeders or other such wetted instruments are provided at halfway or end sections of pipes for city water, feedwater or hot water. These wetted instruments are almost made of copper alloy, such as bronze or brass excellent in castability, mechanical machinability and economical efficiency. Particularly, in valves, joints, etc. made of bronze or brass, alloy added with a prescribed amount of lead is used for bronze ones in order to enhance the characteristics thereof including castability and mechanical machinability and for brass ones in order to enhance the characteristics thereof including cutting machinability and a hot-forging property. When the instruments made of bronze or brass containing lead are exposed to a fluid, such as city water, however, a lead component of the lead containing metal deposited on the wetted surface layer is possibly eluted into the city water. [0003]In view of the above, the water quality standards of city

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water as a beverage have been prescribed by the rating identification method carried out in accordance with the specific procedure. City water has to meet the water quality standards. Since lead is a substance harmful to human bodies and the amount of lead leached has to be reduced as much as possible, restrictions on the lead leaching standard came to strengthening in April, 2003 in Japan. Under these circumstances, wetted instruments of copper alloy produced using a so-called leadless material having lead removed from a fodder and wetted instruments of copper alloy having the elution of lead reduced through surface treatments for a conventional lead-containing material including an acid or alkali cleaning treatment go into circulation. For example, lead elution reduction techniques are described in the following (refer, for example, to Patent Documents 1 to 3).

Patent No. 3345569 (Patent Document 1) was developed by the present applicants and comprises the step of cleaning at least the wetted surface of a piping instrument of copper alloy containing lead with a detergent comprising a nitric acid added with a hydrochloric acid as an inhibitor to form a coat the wetted surface with the hydrochloric acid, thereby deleading the surface layer of the wetted surface. In particular, by using benzotriazole (BTA) in place of the hydrochloric acid as an inhibitor, tarnish and corrosion are suppressed.

[0005] JP-A 2002-180267 (Patent Document 2) discloses, as a lead elution prevention technique, a lead elution prevention treatment method comprising immersing a clasp for feedwater and wastewater in a solution of any of benzotriazole-based compounds to form a film attached firmly to the surface of the clasp.

[0006] JP-A 2001·152369 (Patent Document 3) discloses, as a lead removing technique, a lead elution prevention treatment method comprising immersing a clasp for feedwater and wastewater in an etching

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treatment solution containing an organic carboxylic acid or a salt thereof.

[0007] In addition to the improvement in the prevention of lead elution as described above, an improvement in elution prevention of nickel adversely affecting human bodies has now been calling for urgent attention. Valves, pipe joints, water faucet clasps and other such wetted instruments are subjected to various kinds of plating treatments. commencing with nickel-plating treatment, for the purpose of enhancing the exterior beauty, corrosion resistance and abrasion resistance. Examples ofplating treatments include nickel-plating, nickel-alloy-plating, nickel-chromium-plating and nickel·tin·plating When these nickel-based plating treatments have been adopted, the coat wraps around and adheres to the spout section of a wetted instrument.

[0008] Figure 1 is a cross section showing a JIS wall faucet (made of CAC 406) plated with nickel-chromium alloy, and Figure 2 is a partially enlarged cross section showing a spout section shown in Figure 1. As shown in Figure 2, the spout section of a wetted instrument 1 coated with a coat 2 has a nickel coat 2b not constituting a composite layer with a chromium coat 2a. This is why the nickel coat 2b rather than the chromium coat 2a wraps around the inward portion of the spout section due to the difference in current density range between the two. In this state, when a fluid, such as city water, is supplied to the wetted instrument 1, there is a possibility of nickel of the nickel coat 2b being eluted into the fluid.

[0009] Figure 3 is an enlarged view showing a section "B" in Figure 2 and, as shown, when copper (wetted portion 1a) that is a metal having a high corrosion potential and nickel (nickel coat 2b) that is a metal having a low corrosion potential in contact with each other are wetted with a fluid exhibiting good electrical conductivity, such as city water, to form an electrically conductive state, the nickel (nickel coat 2b)

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having a low corrosion potential is anode-polarized by the copper (wetted portion 1a) having a high corrosion potential to give rise to a corrosion reaction and generate bimetallic corrosion that promotes oxidization and dissolution of nickel. Furthermore, the nickel coat 2b has a plurality of pinholes 2c and, of the pinholes, there exist those reaching the copper surface constituting the under layer of the nickel coat 2b. In consequence of a fluid exhibiting good electric conductivity, such as city water, having entered the pinholes, bimetallic corrosion is also generated there. Moreover, nickel is also eluted from the nickel coat 2b per se. The nickel elution reduction technique is disclosed in the following (refer, for example, to Patent Document 4 or 5).

[0010] The nickel elution reduction treatment method described in JP-A 2002-155391 (Patent Document 4), for example, comprises the steps of plating a water feeder made of copper or copper alloy with nickel, plating the nickel coat surface with chromium and removing the nickel coat running out the chromium coat. In the nickel-removing step, the water feeder is immersed in an oxidizing chemical, such as a sulfuric acid, for removing the nickel coat alone by dissolution. Thus, the nickel coat that has wrapped around the spout at the nickel-plating treatment can be removed by dissolution.

[0011] Japanese Patent No. 2836987 (Patent Document 5) discloses a technique for preventing the elution of nickel comprising the step of forming a thin film of aliphatic unsaturated carboxylic acid on a nickel coat plated on a ceramic substrate on which electronic parts are mounted.

Patent Document 1: Japanese Patent No. 3345569

Patent Document 2: JP-A 2002-180267

Patent Document 3: JP-A 2001-152369

Patent Document 4: JP-A 2002-155391

30 Patent Document 5: Japanese Patent No. 2836987

Disclosure of the Invention:

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Problems the Invention intends to solve:

[0012] However, the lead leaching tests described in JP-A 2002-180267 (Patent Document 2) and JP-A 2001-15239 (Patent Document 3) with the aim of reducing lead leaching is not in accordance with JIS S3200-7 (1997) "City Water Instrument – Leaching Performance Test Method" and, in addition thereto, the amount of leach liquor into which lead has leached out in an actual measurement is unclear. Thus, the technical effect thereof cannot be confirmed.

[0013]In JP-A 2002-155391 (Patent Document 4) with the aim of reducing nickel elution, the nickel coat running out the chromium coat cannot effectively be removed and, since the nickel coat always remains at the wetted section, the nickel component is eluted from this section electrically conductive via a fluid, such as city water, to generate bimetallic corrosion and from the nickel coat per se. Thus, the nickel elution standards cannot satisfactorily be fulfilled at all. In addition, exfoliation of the nickel coat allows copper that is the bare metal to be exposed, and there is a possibility of lead segregated on the surface layer being eluted. Though the nickel elution test described in the prior art is in accordance with JIS S3200-7 (1997) "City Water Instrument - Leaching Performance Test Method," the amount of leach liquor into which nickel has leached out in an actual measurement is unclear. Thus, the technical effect thereof cannot be judged.

In view of the above, a test was conducted under the same conditions as in JP-A 2002-155391 (Patent Document 4) to confirm the technical effect of nickel removal. Though described later in detail in Example 1, the measurement results are shown in Table 6 showing sample No. 9 (using two test pieces each having an area of 2500 mm². An offset was obtained based on the actual measurement. Though described

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later, the offset is a numerical value to be compared with the elution standards. As shown in the same table, the amount of lead eluted was 0.0147 mg/ ℓ and that of nickel eluted was 0.1237 mg/ ℓ . Thus, it was confirmed that the two amounts were too large to satisfy the elution standards, respectively. According to water mixing valves known to have a large amount of nickel elution, therefore, in a sample, the portion of the nickel coat exposed only at the portion that wraps around the spout has an area of 3000 mm². A merely acid-cleaned actual product that has a wetted portion having substantially the same area as the test piece has no longer satisfied the strict nickel elution standards.

[0015]The present inventors have further elucidated the causes of lead elution and nickel elution in this kind of field. Figure 4 is a photograph by an EPMA (Electron Probe MicroAnalyzer) showing the nickel distribution on the inner surface of a JIS wall faucet (made of CAC 406) plated with nickel-chromium alloy and having a nominal diameter of 25A and an inner volume of 40 ml. Figure 5 is a photograph showing the lead distribution. Incidentally, reference numeral 3 in Figure 1 denotes a portion analyzed with an EPMA (Electron Probe MicroAnalyzer). measurement by the EPMA (Electron Probe MicroAnalyzer) was conducted with an acceleration voltage of 30 KV and a probe current of 10 nA. As shown in Figures 4 and 5, in the inner surface (CAC 406 surface) la of the sample 1 subjected to nickel-chromium plating, it is confirmed that lead and nickel partially exist at substantially the same position of the measured surface. As is clear from the electron micrograph of Figure 6, the positions at which the two elements exist are in accord with the positions of the crystal grain boundary of a metal surface.

[0016] Figure 7 is an explanatory view showing the state of existence of lead and nickel at the grain boundary of the inner surface of an instrument, such as a water feeder, the outer surface of which is plated with nickel. Valves, pipe joints, strainers, water faucet clasps or other

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water feeders having complicated flow paths are formed of a copper alloy material by sand casting. The casting surface thus cast is bumpy and, at the dent portions thereof, lead 5 moved from a crystal grain boundary 4 to the surface layer at the time of the solidification is segregated. Particularly at the inside surface layer of a water feeder not subjected to surface treatment, the segregation is conspicuous. When plating treatment is performed in this state, the plating solution stays on the lead 5 at the dent portion and dries there and consequently it is conceivable that a nickel salt 6 different from metallic nickel adheres to the lead 5. Since a water feeder, such as a water faucet clasp, has complicated flow paths and since the plating solution staying inside is difficult to eliminate, it is conceivable that adherence of the nickel salt 6 becomes conspicuous. When a fluid, such as city water, is supplied to the water feeder in this state, the lead 5 and nickel salt 6 are eluted.

In the Patent Documents mentioned above and other prior art, this point is not taken into consideration. Even when adopting the technique of JP-A 2002-180267 (Patent Document 2) to form a film of benzotriazole 33 on lead 31 segregated on a crystal grain boundary 30 as shown in Figure 23, it is insufficient to prevent elution of the lead. Also as shown in the same figure, since the film of benzotriazole 33 is not formed on a nickel salt 32 that is not a metal, elution of the nickel salt 32 consequently proceeds and subsequently a large amount of segregated lead 31 under it is eluted. Thus, it is impossible to prevent elution of both nickel and lead.

Here, the following test was conducted to confirm the effect of reducing leaching of lead by benzotriazole. Two test pieces of pure lead (5 mm x 5 mm x 1 mm) were prepared as samples. One of the test pieces was degreased, then immersed in a 1-wt% benzotriazole solution for five minutes and dried. The other test piece was untreated. The two test pieces were tested for leaching of lead based on JIS S3200-7 (no

conditioning and 16-hour immersion). The test results are as shown in

[0019] [Table 1]

Table 1 below.

	Amount of lead leached		
Sample	(Actual measurement mg/l)		
Untreated sample	0.012		
Sample treated with benzotriazole	0.011		

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[0020] The effect of reducing leaching of lead could not be confirmed from the test results. This implies that the mere immersion treatment with the solution containing benzotriazole fails to form a film of benzotriazole on the lead and, therefore, it is said that leaching of lead cannot be suppressed.

[0021] Further here, it was verified whether or not a film was formed on a nickel salt. A test method was performed when pure water was introduced into a beaker in which a nickel salt in a given amount corresponding to supersaturation was contained and when treated liquid containing 0.5 wt% of benzotriazole + a stearic acid + a small amount of oleic acid was introduced into a beaker. Clear supernatant liquids were obtained and measured in terms of the amount of nickel existing therein in a dissolved state. The clear supernatant liquids were diluted at the same rate and analyzed. Incidentally, 10 g of nickel sulfate and 10 g of nickel chloride were prepared as nickel salts. A test was conducted when 10 cc of pure water was introduced into each of the nickel sulfate and nickel chloride, and 10 cc of treated liquid was introduced into each of the nickel sulfate and nickel sulfate and nickel chloride. The test results are shown in Table 2 below.

[0022] [Table 2]

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	Amount of nickel leached (mg/l)		
Nickel salt	Pure water	Treated liquid	
Nickel sulfate	21.4	25.2	
Nickel chloride	25.7	27.0	

[0023] If a film should be formed on a nickel salt, the effect of reducing leaching of nickel ought to be manifested. However, such an effect could not be confirmed from the test results. It is conceivable that the nickel salt is difficult to bond to benzotriazole because it has no free electron.

Therefore, a mere combination of the technique that cannot prevent the elution of a nickel salt having adhered to the inside residual (Patent Document 2, for example) with the technique that cannot satisfy the nickel-leaching standards (Patent Document 4, for example) cannot attain a technique of reducing nickel leaching satisfying the current nickel-leaching standards. In view of the states where lead or a nickel salt adheres to the inside residual and where a nickel coat adhering to the spout section of a wetted instrument is exposed, a simple combination of the surface treatment technique with the film treatment technique cannot solve the problem under discussion at all. Insofar as organic combinations, such as individual degrees of treatments and order of the treatments, a technique of reducing the elution of nickel and further lead cannot be materialized.

[0025] The technique disclosed in Japanese Patent No. 2836987 (Patent Document 5) is a technique plating a nonmetal, such as ceramic, that is a so-called electroless plating different from the technical means of plating a bare metal with a metal. Therefore, the prior art technique cannot be applied without any modification.

[0026] In consideration of the state of affairs described above, the

present inventors have made keen studies to develop the present invention. The object thereof is to provide a nickel elution prevention method for preventing a wetted instrument made of copper alloy from eluting nickel even when being wetted with a fluid, such as city water, provide a protective film formation agent for preventing the elution of nickel and provide a detergent for preventing the elution of nickel, the wetted instrument including valves for city water, feedwater or hot water, pipe joints, strainers, water faucet clasps, pump supplies materials, water meters, water purifiers, water feeders, hot water feeders or other such wetted instruments each made of copper alloy, such as bronze, brass, etc., plated with a material containing nickel, for example.

Means for solving the Problems:

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[0027] To attain the above object, the invention of claim 1 is directed to a method for preventing elution of nickel from a wetted instrument made of copper alloy and plated with a material containing nickel, comprising applying a protective film formation agent to at least a wetted surface of the wetted instrument to form a protective film, thereby suppressing the elution of the nickel.

[0028] The invention of claim 2 relates to the method for preventing elution of nickel from a wetted instrument made of copper alloy, wherein the protective film formation agent contains at least one species selected from the group consisting of benzotriazole, benzotriazole derivatives and organic acids including straight-chain fatty acids.

[0029] The invention of claim 3 relates to the method for preventing elution of nickel from a wetted instrument made of copper alloy, wherein the protective film comprises two layers of or an appropriate composite layer of benzotriazole and organic acids including a carboxylic acid that constitute the protective film formation agent.

30 [0030] The invention of claim 4 relates to the method for

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preventing elution of nickel from a wetted instrument made of copper alloy, further comprising forming a second protective film on a surface of a nickel coat at a wetted section of the wetted instrument using the protective film formation agent, thereby suppressing the elution of the nickel due to bimetallic corrosion via the second protective film.

[0031] The invention of claim 5 relates to the method for preventing elution of nickel from a wetted instrument made of copper alloy, wherein the nickel coat has pinholes and the second protective film is formed in the pinholes using the protective film formation agent so that the copper alloy and nickel are insulated.

[0032] The invention of claim 6 relates to the method for preventing elution of nickel from a wetted instrument made of copper alloy, wherein the protective film formation agent is used to form the protective film on a surface of a nickel coat on a wetted section of the wetted instrument, thereby suppressing via the protective film dissolution of the nickel per se by wetting.

[0033] The invention of claim 7 relates to the method for preventing elution of nickel from a wetted instrument made of copper alloy, wherein the protective film formation agent is applied to at least the wetted surface of the wetted instrument to form a protective film, and further comprising removing by rinsing a nickel salt adhering as a residual to an inside of the wetted instrument.

[0034] The invention of claim 8 relates to the method for preventing elution of nickel from a wetted instrument made of copper alloy, further comprising deleading a surface layer of the wetted section of the wetted instrument.

[0035] The invention of claim 9 relates to the method for preventing elution of nickel from a wetted instrument made of copper alloy, wherein one or both of the nickel salt adhering as a residual to the inside of the wetted instrument and lead segregated on the surface layer of the

wetted section are rinsed with a cleaning fluid containing a nitric acid and having a hydrochloric acid added thereto as an inhibitor.

[0036] The invention of claim 10 is directed to a protective film formation agent for forming a protective film to prevent elution of nickel from a wetted instrument made of copper alloy, comprising at least one species selected from the group consisting of benzotriazole, benzotriazole derivatives and organic acids including a straight-chain fatty acid.

[0037] The invention of claim 11 is directed to a detergent for preventing elution of nickel from the wetted instrument made of copper alloy, that enables the nickel salt adhering as a residual to the inside of the wetted instrument to be removed and metallic nickel on a spout section of the wetted instrument to be suppressed from being eluted.

The protective film formation component is dissolved in an organic solvent containing at least one species selected from the group consisting of glycolethers, alcohols and amines to be adapted for the DiMethyGlyoximel (DMG) test based on the EN12471 standards. As the glycolethers, 3-methyl-3-methoxybutanol and butyl cellosolve can be raised. As the alcohols, benzylalcohol can be cited. As the amines, morphorine, monoethanolamine, triethanolamine, alcanolamines having an isoform like triisopropanylamine, amines having a cycloform like cyclohexylamine and dicyclohexylamine and long-chain alcoholamine can be listed.

[0039] As the organic solvent, a solvent containing 10 wt% of 3-methyl-3-methoxybutanol and 0.03 wt% or more of morphorine, a solvent containing 10 wt% of 3-methyl-3-methoxybutanol and 0.02 wt% or more of monoethanolamine or a solvent containing 10 wt% of 3-methyl-3-methoxybutanol and 0.05 wt% or more of triethanolamine is preferred.

Effects of the Invention:

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[0040] According to the invention of claim 1, it is made possible to provide an environmentally conscious wetted instrument of copper alloy stably used without eluting nickel from it into a fluid, such as city water, supplied even in a state wherein the wetted instrument is plated with nickel to allow the nickel to adhere to a wetted surface, such as a spout, thereof. The wetted instrument of copper alloy includes water supply instruments installed midway in feed-water pipes or other plumbing, such as valves for city water, feedwater or hot water, pipe joints, strainers, etc. and water supply instruments installed at the end in feed-water pipes, such as water faucets, water purifiers, water feeders, hot water feeders, etc.

According to the invention of claim 2, it is made possible to form a protective film firmly adhering closely to the surface of a nickel coat using a protective film formation agent containing at least one species selected from the group consisting of benzotriazole, benzotriazole derivatives and organic acids, such as a straight-chain fatty acid. Thus, it is made possible to provide a method for preventing elution of nickel from a wetted instrument of copper alloy even when being wetted with a fluid, such as city water.

[0042] According to the invention of claim 3, the protective film can comprise two layers of or an appropriate composite layer of benzotriazole and organic acids including a carboxylic acid that constitute the protective film formation agent. Thus, a further impregnable protective film can be materialized to enable the elution of nickel from a member over the protective film to be prevented infallibly.

[0043] According to the invention of claim 4, a protective film is formed on a wetted portion and on the surface of a nickel coat at the wetted portion to prevent electric conduction between the wetted portion (wetted section) and the nickel coat, thereby preventing the elution of the nickel due to bimetallic corrosion without fail.

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[0044] According to the invention of claim 5, since the defective portions, such as pinholes, of the nickel coat are filled with the protective film formation agent, the elution of nickel due to bimetallic corrosion can infallibly be prevented.

According to the invention of claim 6, the formation of the protective film suppresses dissolution of nickel by wetting the nickel coat per se with a fluid, thus securely preventing the elution of nickel without being affected by a water quality factor, such as pH, and physical and chemical parameters, such as changes in fluid flow and in fluid temperature.

[0046]According to the inventions of claims 7 to 9, in a wetted instrument of copper alloy, such as bronze, brass, etc. plated with nickel, lead segregated on the surface layer of the wetted section thereof and nickel salt adhering as a residual to the inside thereof can securely be removed to prevent the elution of the nickel having adhered to the wetted section, such as a spout, thereof. These metals are not eluted even when a fluid, such as city water, has been supplied to the wetted instrument. According to the ministry ordinance of Ministry of Health, Labor and Welfare on the water quality standards, the lead leaching standard value for feedwater instruments (valves etc.) installed midway in a piping is 0.01 mg/\ell and that for feedwater instruments (water faucets etc.) installed in a piping end is 0.007 mg/l as an exception value. According to the water quality management of Ministry of Health, Labor and Welfare, the target setting item value as regards nickel is $0.01 \text{ mg/}\ell$. Therefore, the nickel leaching standard value for feedwater instruments (valves etc.) installed midway in a piping is 0.01 mg/\ell\$ and that for feedwater instruments (water faucets etc.) installed in a piping end is 0.001 mg/l. It is made possible to provide a method for preventing the elution of nickel from a wetted instrument of copper alloy, which satisfies these standards. Particularly, an acid cleaning treatment with a cleaning fluid containing a

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nitric acid and having a hydrochloric acid added thereto as an inhibitor functions to manifest an activation effect and as a pretreatment of the formation of a protective film. Thus, organic combination of the acid cleaning treatment and the protective film formation treatment can be materialized.

[0047] According to the invention of claim 10, it is made possible to form a protective film firmly adhering closely to the wetted portion of a wetted instrument of copper alloy plated with nickel and to the surface of a nickel coat at the wetted section and also possible to fill the defective portions, such as pinholes, of the nickel coat. Therefore, it is possible to provide a protective film formation agent for forming a protective film capable of preventing elution of nickel due to bimetallic corrosion not to mention by wetting the nickel coat per se with a fluid.

According to the invention of claim 11, it is made possible to provide a detergent for preventing elution of nickel from the wetted instrument, which detergent enables the nickel salt adhering as a residual to the inside of the wetted instrument to be removed and metallic nickel on a spout section of the wetted instrument to be suppressed from being eluted.

[0049] Furthermore, according to the present invention, by applying the protective film formation agent dissolved in an organic solvent to playthings, literary tools, accessories, food processors, medical devices, medical products, etc. made of copper alloy, stainless steel, nickel alloy, steel, etc. and plated with nickel etc., it is possible to provide an excellent effect of preventing an onset of a nickel allergy resulting from contact with these products and parts.

Brief Description of the Drawings:

[0050] Figure 1 is a cross section showing a JIS wall faucet (made of CAC 406) plated with nickel-chromium alloy.

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Figure 2 is a partially enlarged cross section showing a spout section "A" shown in Figure 1.

Figure 3 is an enlarged view showing a section "B" in Figure 2.

Figure 4 is a photograph by an EPMA (Electron Probe MicroAnalyzer) showing the nickel distribution on the inner surface of a JIS wall faucet (made of CAC 406) plated with nickel-chromium alloy.

Figure 5 is a photograph by an EPMA (Electron Probe MicroAnalyzer) showing the lead distribution on the inner surface of a JIS wall faucet (made of CAC 406) plated with nickel-chromium alloy.

Figure 6 is a micrograph by an electron microscope showing the inner surface of a JIS wall faucet (made of CAC 406) plated with nickel-chromium alloy.

Figure 7 is an explanatory view showing the state of existence of lead and nickel at the grain boundary of the inner surface of a wetted instrument plated with nickel.

Figure 8 is a flowchart showing one example of the processing steps in a method for preventing the elution of nickel according to the present invention.

Figure 9 is a cross section showing a commercially available single lever-type mixing faucet (made of CAC 406, having an inner volume of about 155 ml and plated with nickel-chromium alloy) used as a sample.

Figure 10 is a cross-sectional explanatory view showing the state of the surface layer of the wetted part having encountered a protective film formation treatment.

Figure 11 is a graph showing the entire qualitative analysis results having detected as sensitivity (kCPS) photoelectron spectra peculiar to elements emitted from a surface (copper plate) in an analysis region using a top surface hyphenated analytical

instrumentation (XPS = X-ray Photoelectron Spectroscopy).

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Figure 12 is a graph showing the entire qualitative analysis results having detected as sensitivity (kCPS) photoelectron spectra peculiar to elements emitted from a surface (pure nickel copper plate) in an analysis region using a top surface hyphenated analytical instrumentation (XPS).

Figure 13 is a graph showing the entire qualitative analysis results having detected as sensitivity (kCPS) photoelectron spectra peculiar to elements emitted from a surface (copper alloy) in an analysis region using a top surface hyphenated analytical instrumentation (XPS).

Figure 14 is a graph showing the entire qualitative analysis results having detected as sensitivity (kCPS) photoelectron spectra peculiar to elements emitted from a surface (nickel) in an analysis region using a top surface hyphenated analytical instrumentation (XPS = X-ray Photoelectron Spectroscopy).

Figure 15 is a graph showing the qualitative analysis results of a surface in an analysis region using a Fourier Transform Infrared Radiation (FT-IR) spectroscopy analyzer.

Figure 16(a) shows a sample having formed a square region to be wetted through masking of four sides of each of the front and rear surfaces thereof, and Figure 16(b) is a perspective view showing a sample having plating removed over a width "a" to expose bare metal material sections that are to be wetted.

Figure 17 is a graph showing the results of a nickel-leaching test.

Figure 18 is a graph showing the results of a leaching test of Ni and Pb shown in Table 6.

Figure 19 is a graph showing the results of a leaching test of Ni and Pb shown in Table 7.

Figure 20 is a graph showing the results of verifying the treatment time necessary for the formation of a protective film satisfying a nickel-leaching amount of 0.001 mg/l.

Figure 21 is a graph showing the results of verifying the effective concentration of a stearic acid.

Figure 22 is a graph showing the results of verifying the effective concentration of an oleic acid.

Figure 23 is an explanatory view showing the state at the crystal grain boundary of the inner surface of a wetted instrument subjected to the conventional lead elution prevention treatment.

Figure 24 is a graph showing the results of analysis by the Ultraviolet Photoelectron Spectroscopy (UPS) and Penning Ionization Electron Spectroscopy (PIES).

Figure 25 is a graph showing the results of analysis by the Ultraviolet Photoelectron Spectroscopy (UPS) and Penning Ionization Electron Spectroscopy (PIES).

Figure 26(a) is an explanatory plan view showing the structure of a protective film, and Figure 26(b) is an explanatory cross section showing the structure of the protective film.

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Explanation of Reference Numerals:

Nickel salt

Explair	ation	reference rumerals.
[0051]	1	Wetted instrument made of copper alloy
	1a	Wetted section (wetted portion, wetted surface)
	2	Plating
	2a	Chrome plating
	2b	Nickel plating
	2c	Pinhole
	5	Lead

30 13 Deleading step

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15 Plating step

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16 Nickel-removing step

18 Protective film formation step

20 Protective film

20a Protective film (benzotriazole)

20b Protective film (organic acid)

Best Mode for carrying out the Invention:

[0052] A nickel elution prevention method according to the present invention applied to a wetted instrument made of bronze or brass will be described as an embodiment with reference to the drawings. The wetted instrument used herein includes feedwater instruments and their parts installed midway in a feedwater pipe or plumbing, such as valves for city water, feedwater or hot water, pipe joints, strainers, etc.; feedwater instruments and their parts installed at the exit of a feedwater pipe, such as water faucets, water purifiers, water feeders, hot water feeders, etc.; and other completed products connected directly to a feedwater pipe or plumbing.

In making a nickel elution prevention treatment in the present invention, wetted instruments made of copper alloy cast and then processed (valve parts in this example) are preferably disposed in an exclusive net container resistant to heat and chemicals so as not to collide against each other to induce traces and flaws of collision. N workpieces in number that are wetted parts, such as bodies, bonnets, etc. are disposed as a one unit in an exclusive container. The one unit is subjected to treatments described below as a unit to enable the variation of treatment in the parts to be decreased and the quality of the parts to be made constant. Incidentally, parts constituting a valve may be disposed in a lump in an exclusive container and then treated.

[0054] The workpieces are disposed so as not to form air pockets

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in which air bubbles are collected, but to remove the air bubbles to the upper side and sideways. Since the wetted instrument has a complicated shape, during the immersion in each treatment vessel, stimuli by shaking and ultrasonic waves are given to completely remove the air bubbles still remaining. It is thus preferred that the whole wetted surface of the wetted instrument come into contact with a cleaning fluid. In the present example, all the steps described later are taken, with the wetted instrument contained in the exclusive container. After the treatment, the wetted instrument is removed from the exclusive container and brought to an assembling step. An acid cleaning treatment described later may be performed in a state of a completed product (a valve in this example) constituted by a plurality of parts cast and then processed.

[0055]Each of the steps of the nickel elution prevention method according to the present invention will next be described. Figure 8 is a flowchart showing one example of the processing steps in a method for preventing the elution of nickel according to the present invention. A degreasing step 10 is for removing cutting oils and rustproof oils during the processing. The degreasing step is important because insufficient degreasing makes it difficult to completely remove lead during an acid cleaning treatment described layer. Incidentally, when the object to be treated (valve parts in this example) is heavily dirty, it is effective to provide a hot-water washing step 9 prior to the degreasing step 10 to Examples of the degreasing step 10 are remove extraneous matter. shown in Table 3 below. Of these, adoption of an alkali chelate detergent is preferred to prevent an adverse affect of a chlorine based organic solvent on the environment and an increase in BOD by an emulsion detergent.

[0056] [Table 3]

Detergent	Temperature	Time	Cleaning conditions	
Chlorine-based	Room		Immersion and	
organic solvent	temperature	5 min	ultrasonic cleaning	
Neutral emulsion	Room		Immersion and	
detergent	temperature	10 min	ultrasonic cleaning	
Alkali emulsion	Room		Immersion and	
detergent	temperature	10 min	ultrasonic cleaning	
Alkali chelate			Immersion and	
detergent	50°C	10 min	0 min shaking	
Jet steam cleaning	•	5 min		

[0057]When the alkali detergent has been used in the degreasing step 10, the detergent is fully washed away in a water-washing step 11 subsequent to the degreasing step 10. A plurality of water-washing vessels are provided and, for example, an end water washing vessel may contain a mixed acid of 7 wt% of an nitric acid and 7 wt% of hydrochloric acid to completely remove by neutralization the alkali detergent component brought into the end vessel by a container. In a neutralization step 12, by managing the pH (hydrogen-ion exponent) in a main vessel provided for neutralization, a minute amount of the alkali component remaining in the water-washing step 11 can be removed reliably. When a deleading step 13 using a cleaning fluid comprising a mixed acid is to be taken after the neutralization step 12, as in this example, deterioration of the acid by neutralization is prevented to effectively promote deleading reliably.

The deleading step 13 will be described. The deleading step 13 adopts the same cleaning fluid as in a nickel-removing step 16 described later, and the wetted instrument is immersed in the cleaning fluid comprising a nitric acid (at a concentration of 0.5 wt% to 0.7 wt%) and a hydrochloric acid (at a concentration of 0.05 wt% to 0.7 wt%) in a treatment vessel to effectively remove the lead deposited on the surface

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layer of the wetted portion. In the case of a material having a large content of lead, such as CAC 406, since the present step 13 is provided prior to a plating step 15, it is effective because the lead segregated on the copper surface layer in the region on which plating is intended to perform can be removed in advance. While the cleaning fluid used in the deleading step 13 of the present example comprises a mixed acid having a nitric acid added with a hydrochloric acid as an inhibitor, it is used in the form of admixing in city water or pure water an acid, such as a nitric acid, or a mixed acid of a nitric acid added with a hydrochloric acid having an effect of an inhibitor. In this case, since the hydrochloric acid ions Cl corrode the lead while forming a uniform film on the copper surface, the lead is corroded, with the surface retained lustrous. At this time, though lead hydrochloride and lead nitrate are formed on the lead portion, since these salts are soluble in the mixed acid, the corrosion goes on.

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[0059]The acids contained in the cleaning fluid will be described. It is generally known that acids corrode (oxidize) lead. Since lead is easy to form an oxide film as a result of reaction thereof with an acid, the corrosion thereof is difficult to go on. However, lower organic acids, such as a nitric acid, a hydrochloric acid and an acetic acid, continuously corrode lead. Of these, the nitric acid (HNO₃) exhibits the highest lead corrosion rate. On the other hand, since the hydrochloric acid (HCl) exhibiting a low lead corrosion rate as compared with the nitric acid exhibits a high intensity of chemical combination with copper, when acid cleaning is performed using a mixed acid with a nitric acid, a copper chloride (CuCl) coat is formed on the surface of a wetted instrument, prior to the formation of a copper oxide (Cu₂O or CuO) coat by a chemical reaction of the nitric acid with copper, to suppress corrosion of the copper by the nitric acid. Thus, the hydrochloric acid functions as a so-called inhibitor. The inclusion of the hydrochloric acid eliminates oxidation of copper on the surface of a wetted instrument and prevents a disadvantage

in discoloring the surface into black to maintain metallic luster.

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[0060] In addition, ultrasonic cleaning or shaking may be performed in a treatment vessel to promote lead corrosion. The stimulatory effect of lead elution by ultrasonic cleaning or shaking of a wetted instrument will be described. The ultrasonic cleaning is to apply ultrasonic waves to the wetted instrument in a cleaning fluid and has an effect of rapidly remove from the wetted instrument various lead compounds produced by the reaction in the cleaning fluid. The shaking is to shake the wetted instrument per se in the cleaning fluid and has an effect of removing lead compounds from the wetted instrument and eliminating air pockets. Particularly, making stir of the cleaning fluid vigorous forms lead compounds to allow lead to be ready to elute. It is better to use the ultrasonic cleaning and the shaking together.

Though the deleading step adopts the same cleaning fluid as in the nickel-removing step, as described above, this is not limitative. A cleaning treatment with an acid other than the mixed acid as described above or an alkali cleaning treatment will suffice. Also, after the plating step and deleading step are taken, nickel may be removed in the nickel-removing step 16 to be described later. In the case of a material having a small lead content, it goes without saying that both lead and nickel can be removed in the nickel-removing step described later, with the deleading step 13 omitted.

[0062] Though the plating step 15 in this example adopts well-known and well-understood electrolytic nickel-chromium plating, this is not limitative. Nickel plating, nickel alloy plating, nickel-tin plating, etc. can optionally be adopted according to the implementation thereof. The plating contemplated by the present invention is not special plating, such as supercritical plating, but plating to which commercially available feedwater instruments, such as water faucets, valves, etc., are subjected.

[0063]The elution of nickel from a wetted instrument having undergone a plating treatment with a material containing nickel will now For example, in nickel-chromium plating that is electroplating, while the wetted instrument is immersed in a liquid for plating, a chromium layer is formed on the outer surface of the wetted instrument facing an electrode, with nickel as a binder. On the other hand, the inner surface (wetted surface etc.) is formed with no plated coat because it does not face the electrode. Of the wetted surface, the spout sections "A" surrounded by dashed lines in Figure 1 have nickel plating adhering thereto. As shown in Figure 2, however, at the spout section of the wetted surface 1a of the wetted instrument 1 plated with the coat 2, the nickel coat 2b not forming a state of a composite together with the chromium coat 2a is present. This is a result of the nickel coat 2b wrapping around the spout farther inward than the chromium coat 2a due to the difference in current density range between the two.

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[0064]As described in the foregoing and as shown in Figure 3, when copper (wetted portion 1a) that is a metal having a high corrosion potential and nickel (nickel coat 2b) that is a metal having a low corrosion potential in contact with each other are wetted with a fluid exhibiting good electrical conductivity, such as city water, to form an electrically conductive state, the nickel (nickel coat 2b) having a low corrosion potential is anode-polarized by the copper (wetted portion 1a) having a high corrosion potential to give rise to a corrosion reaction and generate bimetallic corrosion that promotes oxidization and dissolution of nickel. Furthermore, the nickel coat 2b has a plurality of pinholes 2c and, of the pinholes, there exist those reaching the copper surface constituting the under layer of the nickel coat 2b. In consequence of a fluid exhibiting good electric conductivity, such as city water, having entered the pinholes, bimetallic corrosion is also generated there. Moreover, nickel is also eluted from the wetted nickel coat 2b per se.

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[0065]In consequence of the analysis of a deeper inside of a wetted instrument by an EPMA (Electron Probe MicroAnalyzer), as shown in Figures 4 to 6, the presence of nickel components can be confirmed. The nickel components are not metallic nickel used in the plating treatment, but nickel salts (nickel sulfate, nickel chloride and nickel hydroxide) contained in a plating solution, having still stayed inside the wetted instrument even after the plating treatment, then dried and adhered to the inner surface. Valves, pipe joints, strainers, water faucet clasps or other water feeders having complicated flow paths are formed of a copper alloy material by sand casting. The casting surface thus cast is bumpy and, at the dent portions thereof, lead moved from a crystal grain boundary to the surface layer at the time of the solidification is segregated. Particularly at the inside surface layer of a water feeder not subjected to surface treatment, the segregation is conspicuous. When plating treatment is performed in this state, the plating solution stays on the lead at the dent portion and dries there and consequently it is conceivable that a nickel salt different from metallic nickel adheres to the lead. Since a water feeder, such as a water faucet clasp, has complicated flow paths and since the plating solution staying inside is difficult to eliminate, it is conceivable that adherence of the nickel salt becomes conspicuous. When a fluid, such as city water, is supplied to the water feeder in this state, both nickel salt and lead are eluted. Incidentally, the dent portions are easy to form on a wetted surface of a mixing faucet particularly intricate in shape due to so-called flow lines formed at casting, and nickel salts are likely to adhere to the dent portions.

[0066] The results of a test for specifying a nickel-leaching source conducted will be described. In this test, the ratio between the case where the leaching source was the nickel salt adhering to the inside of a water feed as a residual and the case where the leaching source was a nickel coat at the spout of a water feeder was confirmed. Figure 9 is a

cross section showing the commercially available single lever-type mixing faucet used as a sample (a product made of CAC 406, having an inside volume of about 155 ml and treated with nickel chromium plating). A leaching test was first performed in accordance with JIS S3200.7, with a sample not wetted beforehand with water, to grasp the amount of nickel to be leached from the entire area of a water feeder. A sample having removed by a cutting process the portions around which a nickel coat wrapped was then prepared, and a leaching test was again performed in accordance with JIS S3200-7, with the sample not wetted beforehand with water, to grasp the amount of nickel salts to be leached from the inside of the water feeder to which the nickel salts adhered as residuals. portions subjected to the cutting process are regions A1 to A3 (having a surface area of about 300 mm²) shown in Figure 9. The results of the tests are shown in Table 4 (actual measurement (mg/l) of Ni and Pb leached in the nickel-leaching source-specifying tests. As shown in Table 4, the ratio of the nickel salts adhering as residuals to the inside of the water feeder to the nickel over the entire area of the water feeder (inside residual ratio) was 10 to 29%.

[0067] [Table 4]

1.770

0.424

0.667

0.617

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	Leaching test		Leaching test		
	(before cutting)		(after cutting)		Ni
Sample	Ni	Pb	Ni	Pb	inside residual ratio
1	0.895	0.547	0.180	0.399	20%

0.205

0.121

[0068] A nickel-removing step 16 will next be described. A wetted instrument is immersed in a cleaning fluid containing a nitric acid and having a hydrochloric acid added thereto as an inhibitor, which fluid is contained in a treatment vessel, thereby removing nickel salts having adhered as residuals to the inside of the wetted instrument. The

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0.600

10%

29%

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cleaning fluid comprises a mixed acid of a nitric acid and a hydrochloric acid added as an inhibitor. Of the mixed acid, the nitric acid acts on nickel to effectively remove the nickel salts from the surface layer of the wetted portion in the form of a nickel nitrate. The cleaning fluid acts also on the nickel adhering to the spout section of the water feeder and has a function to activate the nickel surface as a pretreatment of the protective film formation to be described later. As a consequence, bond between the nickel surface and the protective film is strengthened. Therefore, the nickel elution prevention method in this example adopts an organic combination of the acid cleaning treatment and the protective film formation treatment described later. When the concentration of the cleaning fluid, particularly the concentration of the nitric acid, is thin, removal of the nickel salts from the inside of the water feeder is insufficient and activation of the surface of the nickel adhering to the spout section of the water feeder is also insufficient. On the other hand, when the concentration is unduly thick, since it adversely affects the luster of the nickel coat plated on the outer surface of the water feeder that is an object to be treated, the preferable concentrations of the component acids in the acid cleaning treatment of the present embodiment are set, in which that of the nitric acid falls in the range of 0.5 to 7 wt% and that of the hydrochloric acid falls in the range of 0.05 to 0.7 wt%.

In the case of omitting the deleading step 13 including the case where a material having a small content of lead is an object to be treated, both lead and nickel are removed in the present step 16. While the cleaning fluid is a mixed acid comprising a nitric acid and a hydrochloric acid added thereto as an inhibitor, in the case shown in Figure 7, the nitric acid acts first on the nickel to remove the nickel salts from the surface layer of the wetted portion in the form of a nickel nitrate and thereafter acts immediately on the lead under the removed nickel salt

to remove the lead. In this case, therefore, the single acid cleaning treatment can simultaneously remove both lead and nickel. Incidentally, since the cleaning fluid used in the nickel removing step 16 has been described in detail in the deleading step 13, the description thereof is omitted here. In addition, since nickel is a corrosion resistant material relative to an alkali, such as a sodium hydroxide, and a hydrochloric acid, the solutions thereof cannot remove nickel irrespective of the concentrations thereof and temperatures involved.

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The desirable temperature and time in the acid cleaning treatment will be described. The desirable treatment temperature "x" is in the temperature range expressed as $10^{\circ}\text{C} \le x \le 50^{\circ}\text{C}$. An appropriate temperature range is a room-temperature range that is defined by the range of temperatures taken in a state of the cleaning fluid neither cooled nor heated and variable depending on the temperature of a wetted instrument to be treated and on the atmosphere of the treating vessel exterior. A specific range is from 10°C to 30°C, and the optimal temperature is 25°C. The desirable treatment time "y" is in the range expressed as $5 \text{ min} \le y \le 30 \text{ min}$.

[0071] When the treatment temperature exceeds 50°C, air bubbles by boiling begin to be conspicuous in the cleaning fluid, and air pockets are likely to be formed in the wetted instrument to be treated, with the result that surface portions of the wetted instrument not wetted with the cleaning fluid will possibly exist. Also, evaporation of water and acids will become vigorous, the concentration of the cleaning fluid will become difficult to manage, and the evaporation of the acids will deteriorate the treatment task environment. Therefore, it will be necessary to implement countermeasures to acid resistance for the treatment task area and the task workers. On the other hand, when the treatment temperature falls short of 10°C during the cleaning treatment in winter, the introduction of the cooled wetted instrument into the

treating vessel will possibly decrease the temperature of the cleaning fluid to nearly 0°C to freeze the cleaning fluid. Therefore, the present invention contemplates maintaining the temperature of the cleaning fluid at 10°C or more at which there is no possibility of the cleaning fluid being frozen even when wetted instrument are put into commercial cleaning treatment. In addition, when the treatment time exceeds 30 min, the overage will not contribute too much to the enhancement of the deleading efficiency. Furthermore, the time overage is not adequate to the commercial cleaning treatment. On the other hand, if the treatment time falls short of 5 min, this short-time treatment is insufficient for the elution prevention even when the treatment temperature is elevated. For this reason, the present invention contemplates the treatment time of at least 5 min though this is not limitative.

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[0072] The acid cleaning treatment is also applicable to existing wetted instruments. When the existing wetted instrument is a valve, for example, since nonmetallic parts including packing, gaskets, etc. are also immersed in the cleaning fluid, deterioration thereof is conceivable depending on the cleaning time and temperature and the concentration of the cleaning fluid. In this case, parts made of a material resistant to chemicals, such as fluorine rubber, may be used. While the cleaning fluid used in the acid cleaning treatment of this example contains a hydrochloric acid as an inhibitor, an organic acid, such as an acetic acid or a sulfamic acid, may be mixed with a nitric acid into a mixed acid to remove lead and nickel using the mixed acid.

[0073] After the acid cleaning treatment, the nickel and lead-leaching tests were conducted. The test results will be described here. The acid cleaning treatment as the nickel-removing step was performed using the sample used in the nickel-leaching source-specifying tests. As a result, it was confirmed that almost all nickel salts adhering as a residual to the inside of the water feeder was removed. In the acid

cleaning of the leaching tests, a mixed acid of 4 wt% of a nitric acid and 0.4 wt% of a hydrochloric acid was used. The test results are shown in Table 5 (amounts (mg/ℓ) of Ni and Pb leached after the acid cleaning treatment). As shown in the table, the amount of Ni leached (end offset of plumbing instrument) satisfies the leaching standard of 0.001 mg/ ℓ and the amount of Pb leached (end offset) also satisfies the leaching standard of 0.007 mg/ ℓ .

[0074]

[Table 5]

	N	i	Pb		
	Actual		Actual		
	measurement	End offset	Measurement	End offset	
Sample	(a)	(a)*155/1000	(b)	(b)*155/1000	
2	0.006	0.00093	0.009	0.00140	
3	0.005	0.00078	0.010	0.00155	

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[0075]The wetted instrument is washed with water at the water-washing step 17 after the nickel-removing step to rapidly deprive the surface thereof of the cleaning fluid. Subsequently, the protective film formation treatment is performed at the protective film formation step 18. A protective film forming agent used at the protective film formation step 18 contains at least one species selected from the group consisting of benzotriazole, benzotriazole derivatives and organic acids. Examples of the benzotriazole derivatives include tolyltriazole and carboxybenzotriazole. Examples of the organic acids include straight chain saturated fatty acids like a stearic acid belonging to a carboxylic acid and straight chain unsaturated fatty acids like an oleic acid. As specific examples thereof, a decanoic acid, undecanoic acid, dodecanoic acid, tridecanoic acid, tetradecanoic acid, pentadecanoic acid, stearic acid, heptadecanoic acid, hexadecanoic acid. octadecanoic acid and nanodecanoic acid can be raised. In addition, fatty acids (saturated or unsaturated) other than the straight chain fatty acids, carboxylic acids

having a benzene ring and carboxylic acid ester can also be used.

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[0076]In particular, a protective film-forming agent containing benzotriazole, though adopted at the protective film formation step 1 of the example, is preferred. The reason for it lies in an excellent force of bonding with nickel. Benzotriazole is bonded to the surface of a coat, with a benzene ring having a water-shedding effect (water-repellent property) directed outward and a hydrophilic group directed inward (to the coat side). For this reason, the bonding with the nickel coat can be materialized and, therefore, it is made possible to form a protective film bonded firmly to the nickel coat. Since an organic acid is bonded to a nickel coat, with alkane (C_nH_{2n+2}) having a water-repellent property also directed outward and a hydrophilic group also directed inward, intimate bonding thereof with the nickel coat can be acquired. The force of bonding with the coat surface is obtained in the case of an organic acid due to the delocalization of only the hydrophilic group of the molecules constituting the protective film (motion of electrons), whereas in the case of benzotriazole, it is obtained due to the delocalization of the entire molecules including the benzene ring. That is to say, by using a protective film forming agent possessing a molecular structure having the portion of bonding with the nickel coat highly delocalized, intimate bonding can be attained. Therefore, the protective film forming agent containing benzotriazole is preferred in the point of the strengthened bonding with the nickel coat interdependently with the activation by the acid cleaning treatment. Incidentally, organic substances other than heterocyclic compounds including benzotriazole, benzotriazole derivatives and thiazole or the components enumerated above as the organic acids can be adopted as the protective film forming components insofar as they can intimately be bonded to the surface of the nickel coat. The chemical structures of benzotriazole and organic acids are shown below.

[0077] [Chemical 1]

[0078] [Chemical 2]

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$$C_nH_{2n+2}$$
 $C = O$
 $C = O$
 $C = O$

[0079] When preparing the protective film-forming agent, organic solvents, such as glycol ethers, alcohols and amines capable of dissolving the protective film-forming components in water, are used. As the glycol ethers, 3-methyl-3-methoxybutanol and butylcellsolve can be raised. Benzylalcohole can be cited as the alcohols. Amines include morpholine, monoethanolamine, triethanolamine, alcanolamines having an isoform, such as triisopropanolamine, amines having a cycloform, such as cyclohexylamine and dicyclohexylamine, and long-chain alcohol amines.

At least on the wetted surface 1a of the wetted instrument 1 immersed in the protective film forming agent, a protective film 20 is formed as shown in Figure 10. As a result, the protective film 20 is formed on the surface of the nickel coat 2b on the wetted section 2b and, moreover, the protective film-forming agent enters the small pinholes 2c of the nickel coat 2b. Therefore, the wetted portion 1a and the nickel coat 2b attached to the wetted portion 1a are insulated to prevent both nickel leaching by bimetallic corrosion and nickel leaching from the nickel coat 2b per se when being wetted with a solution.

[0081] The protective film based on the protective film forming

agent is formed on the wetted section at the spout of the water feeder and also on the wetted surface inside the water feeder. Therefore, the lead segregated on the surface layer of the wetted portion is removed at the deleading step 13, and the nickel salts formed inside the water feeder are removed at the nickel-removing step 16. When the protective film forming agent is formed preponderantly of benzotriazole alone, the protective film is formed into a monolayer protective film (protective film 20a in Figure 10). When the protective film forming agent is formed preponderantly of benzotriazole plus an organic acid, the protective film is formed into a bilayer protective film (protective films 20a and 20b in Figure 10) or into a composite protective film in a suitable fashion. When the protective film 20 is formed into a bilayer protective film, as shown in Figure 10, the protective film 20b of the organic acid is deposited on the protective film 20a of the benzotriazole, resulting in the firmly attached protective film to enable prevention of nickel leaching.

[0082] The presence and thickness of the protective film were confirmed using an XPS (top surface hyphenated analytical instrumentation) and an FT-IR (Fourier Transform-Infrared Radiation) spectroscopy analyzer. The explanation thereof is given as follows.

Test 1: Confirmation of presence of protective film on copper alloy and nickel coat

(1) Test Purpose

While it is a conventionally well-established concept that a protective film cannot be formed on so-called metallic nickel, the concept is verified.

(2) Sample

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Test pieces of copper plate (purity: 99.99%) and of pure nickel steel plate (purity: 99.98%) were immersed in an aqueous solution of 1 wt% of benzotriazole at a treatment temperature of 50°C for a treatment time of 5 min. The test pieces were prepared as samples

corresponding to parts a and 8 in Figure 10 and having the protective film 20a of benzotriazole formed thereon. The test piece of pure nickel steel plate was the same as sample No. 21 in Table 8 that was the data of the example to be described later. The aqueous solution of benzotriazole used in this test is an aqueous solution containing a solvent for dissolving benzotriazole in water. This is the same as in the verification test in the prior art described earlier and in tests described later.

(3) Measurement Method

A top surface hyphenated analytical instrumentation (XPS) VG Scientific ESCALAB250 having pass energy (resolution) of 100 eV was used as equipment used. The analysis region (about 0.4 mm) on the samples accommodated in a vacuum chamber was irradiated with X-rays (Al monochlomatic X-ray source, 15 kV, 100 W), with the binding energy thereof varied. The texture and chemical bonding state of the analysis region surface were grasped from the detection of a photoelectron spectrum peculiar to each element emitted from the analysis region surface as the sensitivity (kCPS).

[0083] (4) Measurement Results

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As regards copper plate treated by immersion in benzotriazole:

By irradiating X-rays, with the binding energy increased gradually, as shown in Figure 11, the spectrum "C1s" was detected. By further strongly irradiating the X-rays, the spectrum "N1c" was detected at a region of the copper alloy side slightly deeper than the region at which "C1s" was detected. It was confirmed from the detection of "C1s" that C (carbon) of the benzene ring forming a protective film surface of benzotriazole existed. It was also confirmed from the detection of "N1s" that the nitrogen compounds bonding the benzene ring to the copper-based alloy existed. Thus, it was confirmed that the protective film of benzotriazole was formed on the surface of the copper plate.

[0084] As regards pure nickel steel plate treated by immersion in benzotriazole:

By irradiating X-rays, with the binding energy increased gradually, as shown in Figure 12, the spectrum "C1s" was detected. By further strongly irradiating the X-rays, the spectrum "N1c" was detected at a region of the copper alloy side slightly deeper than the region at which "C1s" was detected. Since the sensitivity of each spectrum was weaker than that of the copper plate, it could be inferred that the thickness of the protective film was slightly smaller than that in the case of the copper plate. However, the detection of "C1s" and N1s" at the same positions (depths) as in the case of the copper plate demolished the conventionally well-established concept that a protective film could not be formed on metallic nickel and confirmed anew that the protective film of benzotriazole was formed on the surface of nickel.

[0085] Test 2: Confirmation of presence of protective film on copper alloy and nickel coat

(1) Test Purpose

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The formation of a protective film on metallic nickel is confirmed, with a protective film forming agent changed. The thickness of each protective film on the surface of copper alloy and nickel coat is also confirmed.

(2) Sample

A sample of copper alloy (CAC 406) subjected to electrolytic plating with nickel (Ni thickness: 2 to 3 µm), part of which was polished to expose the copper alloy was treated by immersion in a treatment fluid containing a protective film-forming agent comprising 0.5 wt% of benzotriazole, a stearic acid and a small amount of oleic acid at a treatment temperature of 50°C for a treatment time of 5 min. The test piece was prepared as a sample corresponding to parts a and B in Figure 10 and having the protective film 20 of benzotriazole and organic acids

formed thereon. This test piece is the same as sample No. 14 in Table 6 that is the data of the example described later. The treatment fluid in this test is an aqueous solution containing a solvent for dissolving the protective film formation components, such as benzotriazole, in water, 0.7 wt% or more of stearic acid and 0.3 wt% or less of oleic acid exclusive of 0 wt% thereof. This is the same as in the verification test in the prior art described earlier and in tests described later.

(3) Measurement Method

It uses the same top surface hyphenated analytical instrumentation (XPS) as in Test 1.

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(4) Measurement Results

As to copper alloy surface:

By irradiating X-rays, with the binding energy increased gradually, as shown in Figure 13, the spectrum "C1s" was detected at the same position (depth) as that in the case of the test piece of Figure 11 having the protective film of benzotriazole formed thereon. While it was confirmed from this fact that C (carbon) existed on the surface of the copper alloy, no spectrum "N1s" as was confirmed in the test piece of Figure 11 was detected even by further irradiating X-rays strongly. Thus, in this measurement, the presence of the protective film of benzotriazole on the surface of the copper alloy could not be confirmed. In spite of the fact that the formation of the protective film of benzotriazole on the surface of the copper alloy was confirmed in the test of Figure 11, it was inferred from no detection in this test is that the spectrum "C1s" was due to the stearic acid and oleic acid. It was conceivable that since the protective film of the stearic acid and oleic acid was thick, no spectrum "N1s" of the benzotriazole existing on the deeper portion of the copper alloy side than the "C1s" detected region could be detected. The chemical formulae of the stearic acid and oleic acid are shown below.

[0087] Chemical formula of stearic acid: C₁₇H₃₅COOH

Chemical formula of oleic acid: C₁₇H₃₃COOH

[0088] As to nickel surface:

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Irradiation of X-rays, with its intensity changed, resulted in detection of "C1s" and "N1s" shown in Figure 14 at the same positions (depths) as in the test piece of Figure 12 having formed the protective film of benzotriazole. It was confirmed anew from the detection that the protective film of benzotriazole was formed on the metallic nickel. When a spectral response of "C1s" was focused on, here, the value thereof detected was much higher than that of the test piece of Figure 12. It was therefore inferred that the protective film of benzotriazole and a protective film composed of the stearic acid and oleic acid coexisted on the nickel surface. When irradiating X-rays further strongly, a spectrum "Ni2p3" was detected on the side of the copper alloy at a region slightly deeper than the "N1s" detection region. It was confirmed from this fact that the thickness of the protective film composed of benzotriazole or stearic acid and oleic acid and formed on the nickel surface was slightly smaller than that of the protective film of benzotriazole or stearic acid and oleic acid and formed on the copper alloy surface.

[0089] Next, as regards the inference that the protective film composed of stearic acid and oleic acid was formed on the protective film composed of benzotriazole, the presence thereof was confirmed by the infrared spectroscopic analysis.

Test 3: Confirmation of presence of protective film on copper alloy and nickel coat

(1) Test Purpose:

It is verified if a protective film composed of stearic acid and oleic acid exists on the protective film of benzotriazole.

(2) Sample:

It is the same as the test piece immersed in 0.5 wt% of benzotriazole plus stearic acid plus oleic acid.

(3) Measurement Method

The equipment used is Fourier Transform Infrared Radiation (FTIR) spectroscopy analyzer Nicolet Nexus 670 having a resolution of 4 cm⁻¹ and a quantity survey of 64 times. By making use of the property that a molecule existing in an analysis region absorbs infrared rays corresponding to vibrational energy when irradiating the infrared rays (laser beams) on the analysis region of a sample accommodated in a vacuum chamber, with their wavenumbers varied, to detect an infrared absorption spectrum (absorbance) (infrared absorption wavenumber) peculiar to each element, the qualitative analysis of the analysis region surface is executed.

[0090] (4) Measurement Results:

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In re the results of analysis of the protective film formation agent:

As shown in Figure 15(a), the peak waveform of the absorption spectrum in the vicinity of a region "\mathcal{T}" is substantially in conformity with that of benzotriazole, indicating that benzotriazole exists in the protective film formation agent. The peak waveforms of the absorption spectra in the vicinity of regions "\Lambda" and "\mathcal{T}" are in conformity with that of a stearic acid ester simple, indicating that stearic acid exists in the protective film formation agent.

[0091] In re the results of analysis of the nickel surface:

The peak waveforms of the absorption spectra in the vicinity of regions " \checkmark " and " $\overset{\circ}{\mathcal{D}}$ " are in conformity with that of the protective film formation agent, as shown in Figure 15(b). This indicates that a protective film of stearic acid is formed on the nickel surface. Here, since the peak waveform of the absorption spectrum in the vicinity of the region " \checkmark " was that overlapping the peak waveform of the absorption spectrum of an oleic acid simple, it was found that the protective film of oleic acid was also formed on the nickel surface. Incidentally, though the

absorption spectrum of benzotriazole is found in the vicinity of the region "\mathcal{T}", the spectral intensity thereof is weak.

[0092] In re the results of analysis of the copper alloy surface:

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The peak waveform of the absorption spectrum in the vicinity of the region " \ref{p} " is substantially in conformity with that of the protective film formation agent as shown in Figure 15(c). This shows that a protective film of oleic acid is formed on the copper alloy surface. Incidentally, no absorption spectrum corresponding to that of oleic acid found on the nickel surface " \ref{p} " shown in Figure 15(b) could not be detected. Furthermore, the absorption spectrum of benzotriazole dose not clearly emerge, a protective film of benzotriazole is present as described above.

After the protective film formation step 18, as shown in Figure 8, drying is carried out at the drying step 19. The drying step 19 evaporates the water content in the protective film-forming agent to fix a protective film of benzotriazole or organic acid firmly to the surface of the copper alloy or nickel coat. Particularly, an organic combination of the acid-cleaning process and the protective film-forming process manifests a synergistic effect by the cleaning fluid and, as a result, a product or member having a protective film formed thereon at the protective film formation step 18 exhibits an extremely excellent effect of preventing nickel leaching. A container having undergone all the steps is conveyed to an assembly step, and the wetted instrument (valve parts in this example) is removed from the container and then subjected to assembly and inspection.

Also, this example takes into consideration environmental issues and wastewater treatment cost. As described above, while an alkali detergent is adopted at the degreasing step 10 of this example, a mixed acid of a nitric acid (concentration: 0.5 wt% to 7 wt%) and a hydrochloric acid (concentration: 0.05 wt% to 0.7 wt%) is used at the

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deleading step 13 and nickel-removing step 16. That is to say, as shown in Figure 8, the alkali detergent contaminated at the degreasing step 10 and the mixed acid solution containing heavy metal at the deleading and nickel-removing steps can react with each other to attain a neutralization process. Precipitates and supernatants can be removed as solids. The oil content can be separated and be subjected to industrial waste disposal. The harmless neutralized water can be leveraged as industrial water. Furthermore, as shown in Figure 8, the diluted alkaline wastewater discharged from the water washing step 11 after the degreasing step 10 and the diluted acidic wastewater discharged from the water washing step 14 after the deleading step 13 and from the water washing step 17 after the nickel-removing step 16 can be mixed to attain a neutralization process, the precipitates and supernatants can be removed as solids, and the oil content can be separated and subjected to industrial waste disposal. The neutralized water now harmless can be leveraged as industrial water. [0095]Furthermore, the protective film formation process of the present invention is applicable to existing assembled and completed products, such as valves, water faucets, etc., having not undergone a nickel-removing step. In this case, the assembled and completed product is degreased and then subjected to a protective film formation treatment. Since the protective film-forming agent of the present invention has no corrosion behavior and has no possibility of packing or gasket that is nonmetal and is incorporated in a valve, water faucet, etc. being deteriorated. This is why the protective film formation treatment can be made relative to the completed products. Incidentally, while a protective film is formed through immersion of a wetted part or completed product in the protective film forming agent, a spraying treatment may be adopted relative to a wetted part having an internal shape liable to produce air pockets in the case of the immersion treatment.

[0096] It was found from the tests for specifying a nickel-leaching

source described above that in the samples that are the water feeders (mixing faucets), those having a spout as the nickel-leaching source occupied about 70%. In view of this, to further pursue whether leaching of nickel from the spout of the water feeder was based on direct leaching from the metallic nickel in the region of the coat or on the bimetallic corrosion due to the potential differential, samples were subjected to the following leaching test.

(1) Test Purpose:

It was verified whether leaching of nickel from the spout of the water feeder was based on direct leaching from the metallic nickel in the region of the coat or on the bimetallic corrosion due to the potential differential.

(2) Sample:

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A plate member having a base metal material of CAC 406 subjected to electrolytic nickel-plating over the entire surface thereof (coat thickness: 2 to 3 µm) was identified as test piece X1 and that subjected to electroless nickel-plating (coat thickness: about 10 µm) as test piece X2. Test piece X1 had a coat thickness of 2 to 3 µm to reproduce the degree of adhesion of a nickel coat to the spout of a wetted instrument and was produced so as to form in the surface thereof a plurality of minute pinholes as shown in Figure 3. Since, in the electroless nickel-plating, pinholes are more difficult to form than in the electrolytic nickel plating and the adhesion of a nickel coat is better than in the electrolytic nickel plating, test piece X2 was treated as a comparative example. Furthermore, a plate member of pure nickel steel (purity: 99.98%) was prepared as test piece Z. Incidentally, test piece X1 corresponds to sample No. 4 in Table 6 showing the data of the examples to be described later, test piece X2 to sample No. 10 in the same table and test piece Z to sample No. 18 in Table 8. The four sides of each test piece on both front and back surfaces thereof were masked as shown in Figure 16(a) to form a

rectangular wetted region of a constant area to be wetted. This masking of the sides of a test piece eliminates unhomogeneous thickness and state of the nickel coats at these sides. While in the leaching test using commercially available water feeders as samples, errors in product and measurement are readily produced depending on the state of casts and the state of plating application in the individual samples, it is possible to eliminate these errors as much as possible and accurately specify the nickel-leaching source in the leaching test using samples making it possible to allow the rated surfaces to be constant. Two pieces of each test piece were prepared, and the total area to be wetted was set to be 5000 mm². This value exceeds the total wetted area of about 3000 mm² of the spouts of the mixing faucets (regions A1 to A3 in Figure 9).

(3) Test Method:

Based on the part test prescribed under JIS S3200-7, the test piece was immersed in a leaching liquid for 16 hours (without conditioning) and the leaching liquid was analyzed with an ICP (Inductively Coupled Plasma) spectrometry. The amount of the leaching liquid was set to be about 160 ml pursuant to the inside volume of the mixing faucet (about 155 ml).

[0097] (4) Test Results:

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The test results are shown in Figure 17. As was clear from the test results, it was found that leaching of nickel from the spout of the water feeder was based on direct leaching from the metallic nickel in the region of the coat or on the bimetallic corrosion due to the potential differential and that the amount of nickel leached based on the latter was larger than that based on the former.

[0098] The prescription under JIS S3200-7 (revised in 2000) will be described here. This prescription prescribes the method of testing the leaching performance of valves, such as water faucets etc., or city water instruments, such as pipes, pipe joints, etc. and the offset of the analysis

results. This leaching test comprises filling the interior of a city water instrument as a sample with a leaching liquid adjusted beforehand to a liquid containing prescribed components, sealing the interior, leaving the leaching liquid as a sample liquid standing for 16 hours and then analyzing components contained in the resultant leaching liquid.

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[0099] In the city water instrument subjected to the leaching test, the interior thereof before being filled with the leaching liquid is subjected to pretreatments including cleaning with city water and water prescribed under JIS K0050 and, when necessary, conditioning that is a treatment comprising repeating cleaning with the leaching liquid and leaving to stand. These pretreatments are prescribed for every one kind of city water instrument to be subjected to the leaching test, specifically, an end water feeder (a water faucet, for example) or a water feeder provided midway in a service pipe or plumbing (a valve, for example). Of the pretreatments, conditioning may be omitted if the sample liquid clearly satisfies the standards without conducting the conditioning. In the examples to be described later, the conditioning was omitted and, on the basis of this prescription, the sample liquid was analyzed using a valve etc. treated by washing with water three times.

[0100] The analysis result of a sample liquid (amount of each component leached) was rated through offset-calculation every one water feeder based on this prescription. In a hot-water mixing faucet that is an end water feeder, in an instrument to be rated having an inside volume of a part in contact with drinking water that is 1ℓ or less, an offset is obtained by dividing the amount actually leached by 1ℓ and multiplying the resultant value by the volume of contact of the instrument to be rated with water to be drunk (155 m ℓ in the water faucet in this example). In a valve that is a water feeder provided midway in a plumbing, an offset is obtained by dividing the amount actually leached by 25 based on the prescription.

Example 1:

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[0101] Examples to which the nickel-leaching prevention method of the present invention is applied will be described in detail along with comparative examples. Plate members each having a base metal material of CAC 406 and nickel-plated were used. Test pieces (having the entire surface plated and a wetted area of about 5000 mm²) subjected to three kinds of plating methods, i.e. electrolytic nickel plating (coat thickness of 2 to 3 µm), electrolytic nickel-chromium plating (Ni coat thickness of 10 µm and Cr coat thickness of 1 µm) and electroless nickel plating (Ni coat thickness of 10 µm) were prepared. A wetted surface formation method and test method for each test piece were the same as in the "tests for specifying a source of nickel leached from the spout." The test results are shown in Table 6 and graphed in Figure 18.

[0102]

[Table 6]

				Amount of	Amount of Ni leached mg/ml	g/m²	Amoun	Amount of Pb leached mg/ml	ed mg/ml
	Sample	Plating			Midway			Midway	
	Š.	_	Treating method	Actual	plumbing	End offset	Actual	plumbing	End offset
				measure.	Offset	(a)*160/1000	measure.	offset	(p)*160/1000
				ment (a)	(a)/25		ment (b)	(b)/25	
			Surface treatment: mixed acid [n-acid(0.5						
		Electrolytic	wt%) + h-acid (0.4 wt%)]; Protective film						
	_	Ni plating	formation: [BTA (0.5 wt%) + stracid +	0.005	0.0002	0.0007	0.001	0.0001	0.0002
Present		Ni: 2 to 3	small amount of o-acid]						
Invention		mi	Protective film formation: [BTA (0.5 wt%)						
	8		+ st-acid + small amount of o-acid	0.069	0.0027	0.0110	0.008	0.0003	0.0013
		Electrolytic	Surface treatment: mixed acid						
		Ni-Cr	[n-acid (0.5 wt%) + h-acid (0.4 wt%)];						
	က	plating	Protective film formation: [BTA (0.5 wt%)	0.001	0.0000	0.0001	0.00	0.0003	0.0014
_		Ni: 10 µm	+ st-acid + small amount of o-acid						
		Cr: 0.1 µm							
	4		Reference (untreated)	0.399	0.0160	0.0638	0.010	0.0004	0.0016
			Surface treatment: mixed acid						
	20	Electrolytic	[n-acid (4 wt%) + h-acid (0.4 wt%)]	0.118	0.0047	0.0189	0.004	0.0002	9000.0
Com.	9	Ni plating	Protective film formation: p-acid (0.9	0.501	0.0201	0.0802	0.007	0.0003	0.0011
Ex.		Ni: 2 to 3							
	2	E E	Protective film formation: p-acid (0.9						
			wt%);	0.471	0.0188	0.0753	0.007	0.0003	0.0012
			Drying at 100°C for 10 min						
_	œ		Surface treatment: mixed acid [n-acid (4						
			wt%) + h-acid (0.4 wt%)]; Protective film	0.099	0.0039	0.0158	0.012	0.0005	0.0019
			formation: p-acid (0.9 wt%)						
	6		Surface treatment: [suracid (200 g/t) +						
			hydrogen peroxide solution (20 g/t)]	0.773	0.0309	0.1237	0.092	0.0037	0.0147
	10	Electroless							
		Ni plating	Reference (untreated)	0.271	0.0108	0.0433	0.020	0.0008	0.0032
		Ni: 10 µm							
min limit	min amid. h.	and bud-pakt	m. and mitting and Theorit' had not been land and BTA: howart more of a cond; at some	- Pina					

n-acid: nitric acid; h-acid: hydrochloric acid; BTA: benzotriazole; stracid: stearic acid; oracid: oleic acid; p-acid: phosphoric acid; su-acid: sulfuric acid

The amount of nickel leached from sample Nos. 1 and 3 having undergone the acid cleaning treatment and the protective film formation treatment with benzotriazole + stearic acid + oleic acid were 0.0007 mg/ ℓ and 0.0001 mg/ ℓ (offsets at the ends in the water feeders), respectively, and satisfied the leaching standard of 0.001 mg/ ℓ at the end in a water feeder. Also, the amount of nickel leached from sample No. 2 having undergone the same protective film formation treatment alone was 0.0027 mg/ ℓ (an offset midway in the plumbing in the water feeder) and satisfied the leaching standard of 0.01 mg/ ℓ midway in the plumbing in a water feeder. Therefore, by performing the surface treatment with the mixed acid of nitric acid + hydrochloric acid before the protective film formation treatment, it was confirmed that the adhesion of the protective film could be made firm and that the nickel-leaching standard of 0.001 mg/ ℓ at the end of a water feeder could be satisfied.

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[0104]On the other hand, in the comparative examples, sample No. 9 having undergone the surface treatment alone with sulfuric acid + hydrogen peroxide solution, and sample Nos. 6 and 7 having undergone the surface treatment alone with phosphoric acid could not satisfy the nickel-leaching standards, and the amounts of nickel leached were rather Incidentally, sample Nos. 5 and 8 having undergone the increased. surface treatment with the mixed acid of nitric acid + hydrochloric acid satisfied the leaching standard midway in the plumbing in a water feeder. [0105]Since the samples were test pieces having lead difficult to segregate on the alloy surfaces, the amount of lead leached from even the samples belonging to the comparative examples (exclusive of sample No. 9) satisfied the lead-leaching standard of 0.01 mg/ ℓ . By subjecting these samples to the treatment according to the present invention, the amount of the lead leached could continuously be suppressed. In sample No. 1, it was made possible to further reduce the amount of the lead leached. Incidentally, in sample Nos. 2, 6 and 7 having undergone the protective

film formation treatment, the amount of the lead leached was not so reduced to a great extent. It could be inferred from this fact that a protective film was not deposited on the lead surface. In sample No. 9, the amount of the lead leached was rather increased.

Example 2:

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Each of plate members having a base material of CAC 406 was subjected to hydrolytic nickel plating (coat thickness: 2 to 3 µm), then masked in the same manner as in the samples in Table 6 and deprived of the coat over a width "a" as shown in Figure 16(b). Test pieces each having the base metal material wetted at the deprived section were prepared. Two pieces of each test piece were prepared, and the ratio of the coat surface was set to occupy about one half the total wetted surface area (about 5000 mm²) (partially plating application sample: coat wetted area of about 2500 mm²). The test method was the same as the "test for specifying the source of nickel leached from the spout." The test results are shown in Table 7 and graphed in Figure 19

[0107]

[Table 7]

				Amoun	Amount of Ni leached mg/ml	ted mg/mf	Amoun	Amount of Pb leached mg/ml	led mg/mf
					Plumbing			Plumbing	
	Sample	Plating	Treating method	Actual	midway	End offset	Actual	midway	End offset
	' N			measure.	offset	(a)*160/1000	measure.	offset	(P)*160/1000
				ment (a)	(a)/25		ment (b)	(b)/25	
Present	11	Electrolytic	Surface treatment: mixed acid [n-acid (4	0.003	0.0001	0.0005	0.023	0.001	0.004
Invention		Ni plating	wt%) + h acid (0.4 wt%)]; Protective film						
	12	Ni: 2 to 3	formation: [BTA (0.5 wt%) + st-acid + small	0.002	0.0002	0.0008	0.011	0.000	0.002
		目	amount of o-acid						
			Surface treatment: mixed acid In-acid (4						
	13		wt%) + h-acid (0.4 wt%)]; Protective film	900.0	0.0002	0.0010	0.053	0.002	0.008
			formation: BTA (1 wt%)						
			Protective film formation: [BTA (0.5 wt%) +						
	14		stracid + small amount of oracid	0.035	0.0014	0.0056	0.103	0.004	0.016
Comp.	15	Electrolytic	Reference (untreated)	0.118	0.0047	0.0189	0.161	9000	0.026
Ä		Ni plating	Surface treatment: mixed acid In acid (4						
	16	Ni: 2 to 3	wt%) + h-acid (0.4 wt%)]	0.236	0.0094	0.0378	0.072	0.003	0.012
		<u> </u>	Surface treatment: [suracid (200 g/l) +						
	17		hydrogen peroxide solution (20 g/k)]	0.291	0.0116	0.0466	0.148	9000	0.024
n-acid: nita	ric acid; h	acid: hydrochlo	n-acid: nitric acid: h-acid: hydrochloric acid: BTA: benzotriazole; st-acid: stearic acid; o-acid: oleic acid; su-acid: sulfuric acid	rid; o-acid: o	leic acid; su	acid: sulfuric a	cid		

[0108]The amount of nickel leached from sample Nos. 11 and 12 having undergone the acid cleaning treatment and the protective film formation treatment with benzotriazole + stearic acid + oleic acid and from sample No. 13 having undergone the protective film formation treatment with benzotriazole were in the range of from 0.0005 mg/l to 0.0010 mg/\(\) (offsets at the ends in the water feeders) and satisfied the leaching standard of 0.001 mg/l at the end in a water feeder. Also, the amount of nickel leached from sample No. 14 having undergone the protective film formation treatment alone with benzotriazole + stearic acid + oleic acid was 0.0014 mg/l (an offset midway in the plumbing in the water feeder) and satisfied the leaching standard of 0.01 mg/\ell midway in the plumbing in a water feeder. Therefore, by performing the surface treatment with the mixed acid of nitric acid + hydrochloric acid before the protective film formation treatment, it was confirmed that the adhesion of the protective film could be made firm and that the nickel-leaching standard of 0.001 mg/l at the end of a water feeder could be satisfied.

On the other hand, in the comparative examples, sample No. 17 having undergone the surface treatment alone with sulfuric acid + hydrogen peroxide solution could not satisfy the nickel-leaching standards. Incidentally, sample No. 16 having undergone the surface treatment with the mixed acid of nitric acid + hydrochloric acid satisfied the leaching standard midway in the plumbing in a water feeder.

[0110] Since the test pieces having the base metal wetted were used as the samples, there was no sample that belongs to the comparative examples satisfying the lead-leaching standard of 0.007 mg/ ℓ . It was confirmed, however, that the samples belonging to the comparative examples when having undergone the treatments of the present invention satisfied the above standard.

Example 3:

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30 [0111] Plate members of pure nickel steel (purity: 99.98%) were

prepared as test pieces (pure nickel steel having a wetted area of about 5000 mm². The wetted surface forming method and test method were the same as the "test for specifying the source of nickel leached from the spout." The test results are shown in Table 8

[0112]

[Table 8]

				Amount	Amount of Ni leached mg/l	mg/ℓ
		Plating			Plumbing	
	Sample No. method	method	Treating method	Actual	Midway	End offset
				Measurement (a)	Offset (a)/25	(a)*(160/1000)
	18		Reference (untreated)	0.029	0.0012	0.0046
			Surface treatment: mixed acid			
	19		[nitric acid (4 wt%)	0.365	0.0146	0.0584
			+ hydrochloric acid (0.4 wt%)]			
			Protective film formation: [BTA			
	20	Pure	(0.5 wt%) + stearic acid	0.020	0.0008	0.0032
Comp.		nickel	+ small amount of oleic acid			
Ex.		steel	Protective film formation:			
	21	(purity:	BTA (1 wt%)	0.021	0.0008	0.0034
		(%8.66	Surface treatment: mixed acid			
-			[nitric acid (4 wt%)			
	22		+ hydrochloric acid (0.4 wt%)]	0.010	0.0004	0.0016
			Protective film formation: [BTA			
			(0.5 wt%) + stearic acid			
			+ small amount of oleic acid			

[0113] While sample Nos. 20 to 22 satisfied the leaching standard midway in the plumbing of a water feeder that was 0.01 mg/ ℓ , they could not satisfy the leaching standard at the end of a water feeder that was 0.001 mg/ ℓ . Also, as shown in the table, the amount of Ni leached from sample No. 19 having encountered the surface treatment with the mixed acid of nitric acid + hydrochloric acid exceeded that from sample No. 18 (reference). This was caused by the fact that the nickel surface was activated with the acid to render the nickel to be easy to bond to the water molecules rather than the fact that the nickel (so-called metallic nickel) was dissolved with the acid. However, this does not mean that the acid corrodes the nickel surface. The following test was conducted to exemplify it.

In this test, three square plate members of pure nickel steel (5 mm x 5 mm x 0.8 mm) were prepared as test pieces. A degreasing agent (organic solvent) was used to degrease the surfaces of the test pieces and then a precision electronic scale (measurable up to a unit of 0.1 mg) was used to the weights thereof. Subsequently, the test pieces were immersed for 10 min in a mixed acid solution comprising 4 wt% of nitric acid + 0.4 wt% of hydrochloric acid, then dried and weighed with the precision electronic scale to confirm the change the weights of the test pieces before and after the treatment with the mixed acid. The measurement results are shown in Table 9.

[0115] [Table 9]

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Test piece	Weight before treatment	Weight after treatment
·	with mixed acid (mg)	With mixed acid (mg)
1	144.3	144.3
2	134.3	134.3
3	143.0	143.0

[0116] As was clear from the table, no change in weight of the test pieces was found before and after the acid-cleaning treatment. Thus, it could be confirmed that the mixed acid solution did not corrode the nickel.

Whit it is conceivable to increase the concentration of the mixed acid and add other acid components in order to enable the mixed acid treatment alone to satisfy the leaching standard for nickel from the end instrument, when a water faucet is actually immersed in such a treatment solution, the coat on the outside surface as well as nickel on the inside surface thereof will possibly separate or exfoliate. According to the treatment of the present invention, it is made possible to suppress the elution of nickel or lead on the inside surface to a level satisfying the prescribed leaching standards without adversely affecting the coat on the outside surface while using the immersing treatment exhibiting good productivity.

Example 4:

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[0117]Next, examples in which the nickel elution prevention method of the present invention is applied to actual products of ball valves will be described in detail together with a comparative example. 10K ball valves (made of C3771) having a nominal diameter of 1/2B, an inside volume of 25 ml and a wetted area, including the inner surface of the valve body and the surface of the ball valve body as the entire inner surface area, of 3440 mm² were used as samples. Each sample was plated with nickel-chromium alloy and subjected to acid-cleaning treatment as the nickel-removing step. The acid-cleaning treatment used a mixed acid of 4 wt% of nitric acid + 0.4 wt% of hydrochloric acid and adopted an immersion treatment of the entire sample at a temperature of 25°C for a period of 10 min to remove nickel and lead as well. In the protective film formation step after the acid-cleaning treatment, a protective film was formed over the entire inside surface of each sample. The treatment conditions adopted a mixture of 0.5 wt% of benzotriazole + stearic acid + a small amount of oleic acid and the immersion treatment of the entire sample at a treatment temperature of 50°C for a treatment period of 5 min. A leaching test was performed without use of conditioning with respect to a water feeder (ball valve)

installed midway in the plumbing in accordance with the prescription of JIS S3200-7 "Leaching Test Method in City Water Instrument." The test results are shown in Table 10.

[0118]

[Table 10]

				Amount of A	Amount of Ni leached mg/l Amount of Pb leached mg/l	Amount of F	b leached mg/l
				Actual	Plumbing	Actual	Plumbing
	Sample	Sample Plating method	Treating method	measure.	midway	measure.	midway
	No.			ment (a)	offset (a)/25	ment (b)	offset (b)/25
Present	23	Electrolytic	Surface treatment: mixed acid	0.201	800'0	0.001	0.000
п	24	nickel-chromium	[nitric acid (4 wt%) +	290.0	0.003	0.055	0.002
		plating	hydrochloric acid (0.4 wt%)]				
			Protective film formation: [BTA				
			(0.5 wt%) + stearic acid + small				
_			amount of oleic acid				
	25		Surface treatment: mixed acid	0.012	0.000	0.025	0.001
			[nitric acid (4 wt%) +				
			hydrochloric acid (0.4 wt%)]				
			Protective film formation: [BTA				
			(1 wt%) + stearic acid + small				
			amount of oleic acid				
	56		Protective film formation: BTA	0.20	0.008	0.100	0.004
			(1 wt%)				•
		Electrolytic	Reference				
Comp. Ex. 27		nickel-chromium	(untreated)	0.400	0.016	0.150	900.0
		plating					

While the amount of nickel leached from sample No. 27 that was a reference (untreated) was 0.016 mg/ ℓ (offset), the amounts of nickel leached from sample Nos. 23 to 25 having undergone the acid-cleaning treatment and the protective film formation treatment with benzotriazole + stearic acid + a small amount of oleic acid and from sample No. 26 having undergone the protective film formation treatment alone with benzotriazole were in the range of 0.000 to 0.008 mg/ ℓ (offset) that satisfied the leaching standard of 0.01 mg/ ℓ midway in the plumbing of a water feeder.

Though the amount of nickel leached from sample No. 27 that was a reference (untreated) was 0.016 mg/ ℓ (offset) that also satisfied the leaching standard of 0.01 mg/ ℓ midway in the plumbing of a water feeder, in sample Nos. 23 to 25 having undergone the acid-cleaning treatment and the protective film formation treatment with benzotriazole + stearic acid + a small amount of oleic acid and sample No. 26 having undergone the protective film formation treatment alone with benzotriazole, the amounts of lead leached were also reduced.

Example 5:

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An example in which the nickel elution prevention method of the present invention is applied to an actual product of water faucet will next be described in detail together with a comparative example. The commercially available single lever-type mixing faucet (made of CAC 406, refer to Figure 9) having an inner volume of about 155 ml and a wetted area of about 3000 mm² as an inner surface area of the sections around which the nickel wrapped (regions A1 to A3 in Figure 9) was used as a sample. After the nickel-plating treatment, the acid-cleaning treatment was performed as the nickel-removing step. The acid-cleaning treatment was an immersion treatment of the entire sample in a mixture of 4 wt% of nitric acid and 0.4 wt% of hydrochloric acid at a temperature of 25°C for 10 min to remove nickel and lead as well. In the protective film

formation step after the acid-cleaning treatment, a protective film was formed over the entire inner surface of the sample. This step was an immersion treatment of the entire sample in a mixture of 0.5 wt% of benzotriazole + stearic acid + a small amount of oleic acid at 50°C for 5 min. A leaching test was conducted without performing conditioning in accordance with the prescription of JIS 3200-7 "Leaching Performance Test Method for City Water Instrument" to measure the leaching amount at the end of a water feeder. The measurement results (single lever-type mixing faucet) are shown in Table 11.

[0122]

[Table 11]

				-				,
				Amo	Amount of Ni	Amo	Amount of Pb	
				leac	leached mg/ℓ	leac	leached mg/l	
-	Sample	Plating	Treatment method	Actual		Actual		
_ -	Š.	method		measure-	End offset	measure-	End offset	
				ment (a)	ment (a) (a)*(156/1000)	ment (b)	ment (b) (b)*(156/1000)	
			Surface treatment: mixed acid					
Present		Electrolytic	[nitric acid (4 wt%) +					
invention	28	nickel-	hydrochloric acid (0.4 wt%)]	900.0	0.000	0.019	0.0029	
		chromium	Protective film formation:					
		plating	[BTA (1 wt%) + stearic acid +	-				
			small amount of oleic acid					
		Electrolytic						
Comp. Ex.	53	nickel-	Reference (untreated)	0.065	0.0101	0.225	0.0349	
		chromium plating						

[0123] While the amount of nickel leached from sample No. 29 that was a reference (untreated) was 0.101 mg/ ℓ (offset), the amount of nickel leached from sample No. 28 having undergone the acid-cleaning treatment and the protective film formation treatment with benzotriazole + stearic acid + a small amount of oleic acid was 0.0009 mg/ ℓ (offset) that satisfied the leaching standard of 0.001 mg/ ℓ at the end of a water feeder.

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The amount of lead leached from sample No. 29 that was a reference (untreated) was 0.0349 mg/ ℓ (offset), whereas the amount of lead leached from sample No. 28 having undergone the acid-cleaning treatment and the protective film formation treatment with benzotriazole + stearic acid + a small amount of oleic acid was 0.0029 mg/ ℓ (offset) that satisfied the leaching standard of 0.007 mg/ ℓ at the end of a water feeder. According to the nickel elution prevention method of the present invention, therefore, it was confirmed that both the nickel-leaching standard and the lead-leaching standard could be satisfied.

In view of the fact that the actual products in Examples 4 and 5 had a complicated structure and did not have a smooth surface, the treating time for forming the protective film was set to be 5 min. It was decided to verify the treatment time necessary for the formation of a protective film capable of satisfying the nickel-leaching amount of 0.001 mg/ ℓ using the test piece X1 of Example 1 having the plate member whose base metal material was CAC 406 subjected to electrolytic nickel plating (nickel coat thickness of 2 to 3 µm). Consequently, the results shown in Table 13 and Figure 20 were obtained. It was found from the results that the immersion for 30-minutes or more enabled the formation of a good protective film.

[0126] [Table 12]

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		Amount of	Ni leached	mg/ℓ
	Protective	Actual	Plumbing	
Treating	film	measure.	midway	End offset
method	formation	ment (a)	offset	(a)*(160/1000)
	time		(a)/25	
Surface treatment:				
nitric acid (4 wt%)	0 sec	0.118	0.0047	0.0189
+ sulfuric acid				
(0.4 wt%)			,	
Surface treatment:				
nitric acid (4 wt%)	5 sec	0.080	0.0032	0.0128
+ sulfuric acid				
(0.4 wt %)	10 sec	0.040	0.0016	0.0064
Protective film				
formation: BTA (0.5	30 sec	0.005	0.0002	0.0008
wt% + stearid acid				
and a small amount of	5 min	0.005	0.0002	0.0008
oleic acid				

Further, here, the effective concentrations of the stearic acid and oleic acid that were the protective film formation components were verified. The test piece X1 of Example 1 having the plate member whose base metal material was CAC 406 subjected to electrolytic nickel plating (nickel coat thickness of 2 to 3 µm) was used as the sample. The test results are shown in Table 13 and Figures 21 and 22. Furthermore, for the purpose of confirming an actual product, the ball valve of Example 4 was used as the sample. The test results are shown in Table 14.

[0128]

[Table 13]

method method measure Midway (a)*(169/1000 measure Midway) Surface treatment: mixed Acid [Midway (a)*(169/1000 measure Midway) Acid [nitric acid (4 wt%) + hydrochloric acid (4 wt%)] Protective film formation: [BTA (0.5 wt%) + stearic acid (0.1 wt%)] Surface treatment: -do- Protective film formation: [BTA (0.5 wt%) + stearic acid (0.5 wt%)] Surface treatment: -do- Protective film formation: [BTA (0.5 wt%) + stearic acid (0.5 wt%)] Surface treatment: -do- Protective film formation: [BTA (0.5 wt%) + stearic acid (0.7 wt%)] OLO wt%) + stearic acid (0.7 wt%)] Surface treatment: -do- Protective film formation: [BTA (0.5 wt%) + oleic acid (0.1 wt%)] Surface treatment: -do- Protective film formation: [BTA (0.5 wt%) + oleic acid (0.3 wt%)] Surface treatment: -do- Protective film formation: [BTA (0.5 wt%) + oleic acid (0.3 wt%)] Surface treatment: -do- Protective film formation: [BTA (0.5 wt%) + oleic acid (0.3 wt%)] Surface treatment: -do- Protective film formation: [BTA (0.5 wt%) + oleic acid (0.3 wt%)] Surface treatment: -do- Protective film formation: [BTA (0.5 wt%) + oleic acid (0.3 wt%)] Surface treatment: -do- Protective film formation: [BTA (0.5 wt%) + oleic acid (0.3 wt%)] Surface treatment: -do- Protective film formation: [BTA (0.5 wt%) + oleic acid (0.3 wt%)] Surface treatment: -do- Protective film formation: [BTA (0.5 wt%) + oleic acid (0.3 wt%)]	Plating	Treating	Amount of	Amount of nickel leached mg/8	mg/ℓ	Amount of	Amount of Pb leached mg/l	3/8
Surface treatment: mixed Midway Action Midway Midway Midway Measure Midway Meath (a) Midway Meath (b) Ment (c) Ment (c) Ment (c) Ment (c) Ment (c) Measure Meath (c) Measure Meath (c) Measure Meath (c) Meath (c)	method	method	Actual	Plumbing	End offset	Actual	Plumbing	End offset
Surface treatment: mixed		P	measure.	Midway	(a)*(169/1000	measure-	Midway	(b)*(169/1000
Surface treatment: mixed			ment (a)	Offset (a)/25		ment (b)	Offset (b)/25	
Acid [nitric acid (4 wt%) +	Electrolytic	Surface treatment: mixed						-
bydrochloric acid (0.4 wt%)] bydrochloric acid (0.1 wt%)] Co.5 wt%) + stearic acid (0.1 wt%)] Surface treatment: -do Protective film formation: [BTA (0.5 wt%) + stearic acid (0.3 wt%)] Surface treatment: -do Protective film formation: [BTA (0.5 wt%) + stearic acid (0.5 wt%)] Surface treatment: -do Protective film formation: [BTA (0.5 wt%) + stearic acid (0.7 wt%)] Surface treatment: -do Protective film formation: [BTA (0.5 wt%) + stearic acid (0.7 wt%)] Surface treatment: -do Protective film formation: [BTA (0.5 wt%) + oleic acid (0.1 wt%)] Surface treatment: -do Protective film formation: [BTA (0.5 wt%) + oleic acid (0.3 wt%)] Surface treatment: -do Protective film formation: [BTA (0.5 wt%) + oleic acid (0.3 wt%)] Surface treatment: -do Protective film formation: [BTA (0.5 wt%) + oleic acid (0.3 wt%)] Surface treatment: -do Protective film formation: [BTA (0.5 wt%) + oleic acid (0.5 wt%)] Surface treatment: -do Protective film formation: [BTA (0.5 wt%) + oleic acid (0.5 wt%)]	nickel	Acid [nitric acid (4 wt%) +						
0.5 µm Protective film formation: [BTA (0.5 wt%) + stearic acid (0.1 wt%)] 0.15 0.0060 0.024 0.024	plating	hydrochloric acid (0.4 wt%)]		-				
6)] 0.15 0.0060 0.024 0.024 6)] 0.10 0.0040 0.016 0.023 6)] 0.005 0.0020 0.008 0.024 6)] 0.006 0.0002 0.0010 0.025 0.003 0.0001 0.0005 0.023 0.006 0.0002 0.0010 0.021	$Ni=2$ to 3 μ m	Protective film formation: [BTA						
6)] 0.10 0.0040 0.016 0.023 6)] 0.005 0.0020 0.008 0.024 6)] 0.006 0.0002 0.0010 0.025 0.003 0.0001 0.0005 0.023 0.006 0.0002 0.0016 0.021		(0.5 wt%) + stearic acid (0.1 wt%)	0.15	0900.0	0.024	0.024	0.001	0.0038
6)] 0.10 0.0040 0.016 0.023 6)] 0.05 0.0020 0.008 0.024 6)] 0.06 0.0002 0.0010 0.025 0.003 0.0001 0.0005 0.023 0.006 0.0002 0.0016 0.021		Surface treatment: -do-						
6)] 0.10 0.0040 0.016 0.023 6)] 0.05 0.0020 0.008 0.024 6)] 0.006 0.0002 0.0010 0.025 0.003 0.0001 0.0005 0.023 0.006 0.0002 0.0016 0.021		Protective film formation: [BTA						
6)] 0.06 0.0020 0.008 0.024 6)] 0.006 0.0002 0.0010 0.025 0.003 0.0001 0.0005 0.023 0.006 0.0002 0.0016 0.021		(0.5 wt%) + stearic acid (0.3 wt%)	0.10	0.0040	0.016	0.023	0.001	0.0037
6)] 0.005 0.0020 0.008 0.024 6)] 0.006 0.0002 0.0010 0.025 0.003 0.0001 0.0005 0.023 0.006 0.0002 0.0016 0.021		Surface treatment: -do-						
6)] 0.05 0.0020 0.008 0.024 6)] 0.006 0.0002 0.0010 0.025 0.003 0.0001 0.0005 0.023 0.006 0.0002 0.0016 0.021		Protective film formation: [BTA						
6)] 0.006 0.0002 0.0010 0.025 0.003 0.0001 0.0005 0.023 0.006 0.0002 0.0010 0.021		(0.5 wt%) + stearic acid (0.5 wt%)]	0.05	0.0020	0.008	0.024	0.001	0.0038
6)] 0.006 0.0002 0.0010 0.025 0.003 0.0001 0.0005 0.023 0.006 0.0002 0.0010 0.021		Surface treatment: -do-						
6) 0.006 0.0002 0.0010 0.025 0.003 0.0001 0.0005 0.023 0.006 0.0002 0.0010 0.021		Protective film formation: [BTA						
0.003 0.0001 0.0005 0.023 0.006 0.0002 0.0010 0.021 0.01 0.0016 0.022		(0.5 wt%) + stearic acid (0.7 wt%)	0.006	0.0002	0.0010	0.025	0.001	0.0040
0.003 0.0001 0.0005 0.023 0.006 0.0002 0.0010 0.021 0.01 0.0016 0.022		Surface treatment: -do-						
0.003 0.0001 0.0005 0.023 0.006 0.0002 0.0010 0.021		Protective film formation: [BTA						
0.006 0.0002 0.0010 0.021		(0.5 wt%) + oleic acid (0.1 wt%)]	0.003	0.0001	0.0005	0.023	0.000	0.0037
0.006 0.0002 0.0010 0.021		Surface treatment: -do-						
0.006 0.0002 0.0010 0.021		Protective film formation: [BTA						
n: [BTA 0.01 0.0004 0.0016 0.022		(0.5 wt%) + oleic acid (0.3 wt%)]	900'0	0.0002	0.0010	0.021	6000.0	0.0034
0.01 0.0004 0.0016 0.022		Surface treatment: -do-						
0.01 0.004 0.002 0.002 0.002 0.0016 0.002		Protective film formation: [BTA						
מכינת (סיס אוניים) סיסים סיסים		(0.5 wt%) + oleic acid (0.5 wt%)]	0.01	0.0004	0.0016	0.022	0.0009	0.0035

[0129]

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[Table 14]

		Amo	ount of Ni lead	ched mg/l
Plating	Treatment	Actual	Plumbing	
method	method	measure-	midway	End offset
		ment (a)	offset $(a)/25$	(a)*(160/1000)
Electrolytic nickel- chromium plating	Surface treatment: mixed acid [nitric acid (4 wt%) + hydrochloric acid (0.4 wt%)] Protective film formation: [BTA (0.5 wt%) + oleic acid (0.1 wt%)]	0.004	0.00016	0.0006

[0130] It was found from these test results that the effects could be manifested when the content of the stearic acid was 0.7 wt% or more and when the content of the oleic acid was 0.3 wt% or less. It was exemplified that these effective contents were applicable to the actual products.

Since C3771 is at a disadvantage in that it induces corrosion by dezinfication, the copper-based alloy developed by the present applicant (JP-A HEI 7-207387) is used to enable a wetted instrument having dezincing resistance and dezincing resistance characteristics to be provided. The copper-based alloy has a composition comprising 59.0 to 62.0% of Cu, 0.5 to 4.5% of Pb, 0.05 to 0.25% of P, 0.5 to 2.0% of Sn, 0.05 to 0.30% of Ni and the balance of Zn and unavoidable impurities (all in wt%) and is excellent in corrosion resistance and hot working. An alternative is a copper-based alloy that has a composition comprising 59.0 to 62.0% of Cu, 0.5 to 4.5% of Pb, 0.05 to 025% of P, 0.5 to 2.0% of Sn, 0.05 to 0.30% of Ni, 0.02 to 0.15% of Ti and the balance of Zn and unavoidable impurities (all in wt%) and having an α + β texture uniformly compartmentalized and is excellent in corrosion resistance and hot working.

[0132] Furthermore, use of the copper-based alloy developed by

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the present applicant (JP-A HEI 10-317078) enables the provision of a wetted instrument excellent in hot working and stress corrosion cracking resistance characteristics besides the aforementioned characteristics. The copper based alloy has a composition comprising 58.0 to 63.0% of Cu, 0.5 to 4.5% of Pb, 0.05 to 0.25% of P, 0.5 to 3.0T of Sn, 0.05 to 0.30% of Ni and the balance of Zn and unavoidable impurities (all in wt%) and having an $\alpha + \beta$ texture uniformly compartmentalized and is excellent in corrosion resistance and hot working. This alloy, when having further undergone an appropriate drawing work and heat treatment, is enhanced in mechanical properties including tensile strength, proof stress and elongation and, when having been satisfactorily deprived of internal stress, excels in stress corrosion cracking resistance. An alternative is a copper-based alloy having a composition comprising 58.0 to 63.0% of Cu. 0.5 to 4.5% of Pb, 0.05 to 0.25% of P, 0.5 to 3.0% of Sn, 0.05 to 0.30% of Ni, 0.02 to 0115% of Ti and the balance of Zn and unavoidable impurities (all in wt%) and having an $\alpha + \beta$ texture uniformly compartmentalized to exhibit excellent corrosion resistance and excellent hot working. This alternative, when having further undergone an appropriate drawing work and heat treatment, is enhanced in mechanical properties including tensile strength, proof stress and elongation and, when having been satisfactorily deprived of internal stress, excels in stress corrosion cracking resistance. It is preferred to have a composition ratio of P and Sn satisfying P (%) x 10 = (2.8 to 3.98 (%)) - Sn (%).

[0133] When using the copper-based alloy developed by the present applicant (JP-A 2000-319736, it is possible to provide a Pb-free copper-based alloy excellent in dezincing resistance, cutting processing property and hot forgeability. The copper-based alloy is characterized in that it contains a component that has a boiling point lower than that of a base phase comprising an α -phase, ($\alpha + \beta$)-phase or ($\alpha + \beta + \gamma$)-phase and a component enabling the base phase and low boiling-point component to be

dispersed to cause uniform dispersion with a soft phase (Bi) and a hard phase (Se + Cu, Se + Zn, γ-phase), thereby enhancing the cutting processing property thereof. The brass for hot forging has a composition comprising 59.0 to 62.0% of Cu, 0.5 to 1.5% of Sn, 1.0 to 2.0% of Bi, 0.03 to 0.20% of Se, 0.05 to 0.20% of Fe and 0.05 to 0.10% of P. The brass for cutting processing is a lead-free alloy that has a composition comprising 61.0 to 63.0% of Cu, 0.3 to 0.7% of Sn, 1.5 to 2.5% of Bi, 0.03 to 0.20% of Se, 0.1 to 0.30% of Fe and 0.05 to 0.10% of P.

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[0134] When using the copper-based alloy developed by the present applicant (JP-A 2003-92217), it is possible to provide a Pb-free copper-based alloy improved in debasement of toughness at high temperature and making the mechanical properties close further to those of CAC 406. The alloy contains at least 2.8 to 5.0 wt% of Sn, 0.4 to 3.0 wt% of Bi and $0 < Se \le 0.35$ wt% to enhance the mechanical properties while securing the prescribed cutting machinability and casting soundness. To be specific, it is a lead-free copper-based alloy having a composition comprising 0.4 to 3.0 wt% of Bi, $0 < Se \le 0.35$ wt%, 2.8 to 5.0 wt% of Sn, 5.0 to 10 wt% of Zn, 3.0 wt% or less of Ni, less than 0.5 wt% of P and less than 0.2 wt% of Pb.

[0135] When subjecting the copper-based alloys represented by the above examples and excellent in corrosion resistance, hot working and stress corrosion cracking resistance characteristics to the nickel elution prevention method of the present invention, it is made possible to provide a wetted instrument made of copper-based alloy having a property of preventing the elution of nickel besides the aforementioned features. Furthermore, use of the lead-free copper-based alloys represented by the above examples enables the provision of copper-based alloys extremely small in the amount of lead eluted. In this case, the neutralizing step 12, deleading step 13 and water washing step 14 can be eliminated from the treatment processing for the method of preventing nickel elution shown in

Figure 8. It is made possible to provide, via the nickel-plating step 15, nickel-removing step 16 and protective film formation step 18, a wetted instrument of copper-based alloy capable of preventing the elution of lead and nickel.

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[0136] Moreover, the protective film formation composition may be dissolved in an organic solvent containing at least one species selected from the group consisting of glycol ethers, alcohols and amines to adapt the result alloy for the DMG test based on the EN 12471 standard. The glycol ethers include 3-methyl-3-methozy butanol and butyl cellosolve, for example. The alcohols include benzyl alcohol. As the amines, morphorine, monoethanolamine, triethanolamine, alcanolamines having an isoform like triisopropanylamine, amines having a cycloform like cyclohexylamine and dicyclohexylamine and long-chain alcoholamine can be raised.

[0137]The DMG test based on the EN 12471 standard comprises a pretreatment of corroding the surface of alloy with artificial sweat imitating the effect of sweat, heat treatment (50°C) and subsequent reaction of nickel ions with dimethylglyoxym in the presence of ammonium into a red color. A cotton-tipped swab is infiltrated with these referential substances and used to wear away part of a test object and observe a change in color at that part. This test is a strict one taking into consideration the inference that anyone would directly touch the nickel-containing substance. Slight tarnish is regarded as having exceeded 0.0005 mg/cm²/week. No tarnish can only be regarded as having passed the test. Test piece X1 that was the plate member whose base metal material was CAC 406 and which underwent electrolytic nickel plating over the entire surface thereof (nickel coat thickness of 2 to 3 µm was used as in Example 1, subjected to the surface treatment with a mixed acid of nitric acid (4 wt%) + hydrochloric acid (0.4 wt%) and then to nickel elution prevention treatment with the protective film formation

agents shown in Table 15 and further subjected to the DMG test under the EN 12471 standard and the leaching test under JIS 3200-7. The results of the tests are shown in Table 15.

[0138]

[Table 15]

	:				Ιſ	Protective	film formati	Protective film formation agent (wt%)	(%)	;	9,	
		No. 1	No. 2	No. 3	No. 4	No. 5	No. 6	No. 7	No. 8	No. 9	No. 10	No. 11
Protective film	Benzotriazole	0.5	0.6	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5
formation										,	,	,
component	Oleic acid	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1
	3-methyl-3-											,
	methoxybutanol	33.	20	10	10	10	10	10	2	10	10	01
	Morpholine	0.02	0.02	0.02	0.03	0.40	•	-				
	Monoethanolamine				•		•	•	0.02	0.28		
Solvent	Triethanolamine						•				0.05	0.67
	Ethanol	33			•	•	•	•		•	-	
	NaOH				•	•	0.01	0.18		•	•	
	Softened water	Balance	Balance	Balance	Balance	Balance	Balance	Balance	Balance	Balance	Balance	Balance
Leaching test ur	Leaching test under JIS S3200-7	Pass	Pass	Pass	Pass	Pass	Rejection	Rejection	Pass	Pass	Pass	Pass
DMG test under	DMG test under EN12471 standard	Rejection	Pass	Rejection	Равв	Pass	Rejection	Rejection	Pass	Pass	Pass	Pass

[0139] While No. 1 shown in Table 15 contained 33 wt% of 3-methyl-3-methoxybutanol, 33 wt% of ethanol, etc. as solvents extremely effective for water-solubilizing the protective film formation components comprising benzotriazole and oleic acid, the result of the test was regarded as being rejectable. Since No. 2 used the solvent containing no ethanol from the standpoint of suppressing the volatilization and containing 3-methyl-3-methoxybutanol, the concentration of which was increased to 50 wt%, it passed the test. However, since it contained a large amount of solvent components, the inflammability became high and the strong chemical small was an obstacle to the protective film formation work. In view of this fact, the concentration of 3-methyl-3-methoxybutanol was fixed to be 10 wt% and other solvent components were adjusted to make attempts to improve the drawbacks.

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[0140] As a result, No. 3 to No. 5 adopting the organic solvent containing 10 wt% of 3·methyl·3·methoxybutanol and 0.03 wt% or more of morpholine, No. 8 and No. 9 adopting the organic solvent containing 10 wt% of 3·methyl·3·methoxybutanol and 0.02 wt% or more of monoethanolamine and No. 10 and No. 11 adopting the organic solvent containing 10 wt% of 3·methyl·3·methoxybutanol and 0.05 wt% or more of triethanolamine, in each of which organic solvent the protective film formation components were dissolved, passed the two tests.

In spite of the fact that No. 1 and No. 10 have the same protective film formation composition, No. 1 failed to pass the DMG test. In order to verify the reason for it, the CASS test under JIS Z2371 was conducted. The corrosion mechanism lies in addition to the effects of corrosion by hydrogen ions resulting from the reduction in pH and corrosion by copper ions accompanied by copper chloride besides corrosion by salt water, such as by salt water spraying. Furthermore, the treatment temperature is set at 50°C to heighten the corrosion speed. Also, though pure Ni steel is to be used by nature, since no discoloration

(rust) is produced even on the corroded surface of the pure Ni steel, no discrimination is possible. Therefore, when pure copper was used as an alternative, almost no discernible corrosion could be found in the case of using the protective film formation agent of No. 10, whereas the film exfoliated and the entire surface was corroded in the case of using the protective film formation agent of No. 1.

Therefore, by specifying the protective film formation components and the solvents for solving the components in water, it is made possible to form a film capable of passing the DMG test under the EN 12471 standard on the nickel-plated surface while suppressing the effects of inflammability and chemical smell during the course of the protective film formation work. Thus, it is possible to prevent nickel allergosis resulting from contact with copper-based alloy plated with nickel.

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Test pieces of pure Ni steel to which No. 1 and No. 10 in Table 15 was applied were analyzed with respect to the states of organic thin films using the Ultraviolet Photoelectron Spectroscopy (UPS) and the Penning Ionization Electron Spectroscopy (PIES). The UPS and PIES analyses are means for observing an organic thin film by making the beams as small as possible and irradiating the beams in parallel to the objective workpieces.

[0144] Figures 24 and 25 show the results of analyses by the Ultraviolet Photoelectron Spectroscopy (UPS) and the Penning Ionization Electron Spectroscopy (PIES). It is found from the analysis results of the protective film formation agent No. 1 shown in Figure 24 that the peak of phenyl group exists in the benzotriazole molecule. It is conceivable in consideration of the results of the molecular crystal growth and CASS test that the crystal growth is localized and that the film structure is unhomogeneous. It is inferred that light beams in the PS and PIES collide against the protuberant benzotriazole crystal mass to seize the

peak of the phenyl group.

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On the other hand, from the results of the analysis of the protective film formation agent No. 10 shown in Figure 25, the oleic acid molecules witheringly small in number seize the peaks of the existing alkyl group and C·O bond. Also in consideration of the CASS test results, it is conceivable that the film has a homogeneous structure and it is inferred that the UPS and PIES beams collide on the tall crystal flux of the oleic acid to seize the peaks of the alkyl group and C·O bond. As shown in Figure 26(a), the arrayed crystals of the oleic acid small in number of molecules are grown in a slender mesh shape like a grid, and the surrounded cell portions are filled regularly with benzotriazole like "go" stones on a "go" board. Moreover, as shown in Figure 26(b), it is conceivable that each crystal is not perpendicular to, but is inclined relative to the analyzed peak.

[0146]In the light of the aforementioned observations, the structure of the protective film will be described in detail in line with the formation process. When using a protective film formation agent comprising benzotriazole, oleic acid, an organic solvent and water, the solvent components first evaporate, and the oleic acid insoluble in water is separated from the water. Subsequently, the oleic acid directs its hydrophilic group toward the nickel side (inside) and its alkyl group (including alkane) having a water repellent property toward the non-nickel side (outside) to form on the nickel coat surface arrayed crystals rising aslant in a mesh shape. The aqueous solution containing benzotriazole is repelled by the arrayed crystals of the oleic acid to enter between the arrays of the oleic acid crystals. The benzotriazole in the aqueous solution has its hydrophilic group positioned along the hydrophilic group of the oleic acid on the nickel side (inside) and its benzene ring that is the water repellent portion thereof positioned along the alkyl group that is the water repellent portion of the oleic acid on the

outside. That is to say, the benzotriazole is filled in the surrounded cells between the arrays of the oleic acid crystals in a rising state in parallel to the oleic acid and, when the water has evaporated, is condensed in the cells to dispose the water repellent portions of the oleic acid and benzotriazole side by side on the outer surface of the protective film.

[0147] A protective film composed of benzotriazole has a structure in which the benzotriazole that is the plane molecule is merely stacked on the nickel coat surface in parallel and readily exfoliates. In a protective film composed of a combination of benzotriazole with an organic acid in the present invention, however, since the benzotriazole is condensed in a state of rising from the coat surface, as described above, the protective film per se is very tough. Of the protective film formation agents shown in Table 15 and having passed the DMG test under the EN 12471 standard, those containing triethanol as a solvent have revealed the best CASS test results and are thus preferred further because it is conceivable that the organic acid (oleic acid) makes the crystal arrays (lattices) dense.

By applying a protective film formation agent having an organic solvent dissolved therein according to the present invention to personal ornaments, such as finger rings, necklaces, pierced earrings, earrings, watches (their straps), eyeglasses (their frames), etc. playthings, such as mini cars, dolls, etc., stationery, such as pencil boxes, clips, etc., medical appliances, such as surgical knives, hype-sticks, etc. welfare and care equipment, such as invalid chairs, axilla clutches, etc., ornamental articles, food-processing equipment, medical products, etc. made of copper alloy, stainless steel, nickel alloy, steel, etc. plated with a material containing nickel, it is made possible to prevent an onset of a nickel allergy resulting from contact with these products and parts.

Industrial Applicability:

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[0149] The nickel elution prevention method, protective film 30 formation agent for prevention of nickel elution and detergent for prevention of nickel elution according to the present invention can be applied to wetted instruments made of copper alloy including bronze and brass and to various kinds of products and parts made of stainless steel, nickel alloy, steel, etc. and can widely be provided in various fields as means for preventing nickel from being eluted with exactitude and lead from being eluted as well.

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